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(54) Title: HAEMOPHILUS ADHERENCE AND PENETRATION PROTEINS

(57) Abstract

Haemophilus adhesion and penetration proteins, nucleic acids, vaccines and monoclonal antibodies are provided.

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### HAEMOPHILUS ADHERENCE AND PENETRATION PROTEINS

### FIELD OF THE INVENTION

The invention relates to *Haemophilus* adhesion and penetration proteins, nucleic acids, and vaccines.

## BACKGROUND OF THE INVENTION

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Most bacterial diseases begin with colonization of a particular mucosal surface (Beachey et al., 1981, J. Infect. Dis. 143:325-345). Successful colonization requires that an organism overcome mechanical cleansing of the mucosal surface and evade the local immune response. The process of colonization is dependent upon specialized microbial factors that promote binding to host cells (Hultgren et al., 1993 Cell, 73:887-901). In some cases the colonizing organism will subsequently enter (invade) these cells and survive intracellularly (Falkow, 1991, Cell 65:1099-1102).

Haemophilus influenzae is a common commensal organism of the human respiratory tract (Kuklinska and Kilian, 1984, Eur. J. Clin. Microbiol. 3:249-252). It is a human-specific organism that normally resides in the human nasopharynx and must colonize this site in order to avoid extinction. This microbe has a number of surface structures capable of promoting attachment to host cells (Guerina et al., 1982, J. Infect. Dis. 146:564; Pichichero et al., 1982, Lancet ii:960-962; St. Geme et al., 1993, Proc. Natl. Acad. Sci. U.S.A.

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90:2875-2879). In addition, H. influenzae has acquired the capacity to enter and survive within these cells (Forsgren et al., 1994, Infect. Immun. 62:673-679; St. Geme and Falkow, 1990, Infect. Immun. 58:4036-4044; St. Geme and Falkow, 1991, Infect. Immun. 59:1325-1333, Immun. 59:3366-3371). As a result, this bacterium is an important cause of both localized respiratory tract and systemic disease (Turk, 1984, J. Med. Microbiol. 18:1-16). Nonencapsulated, non-typable strains account for the majority of local disease (Turk, 1984, supra); in contrast, serotype b strains, which express a capsule composed of a polymer of ribose and ribitol-5-phosphate (PRP), are responsible for over 95% of cases of H. influenzae systemic disease (Turk, 1982, Clinical importance of Haemophilus influenzae, p. 3-9. Sell and P.F. Wright (ed.), In S.H. Haemophilus influenzae epidemiology, immunology, and prevention of disease. Elsevier/North-Holland Publishing Co., New York).

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20 The initial step in the pathogenesis of disease due to influenzae involves colonization of the upper respiratory mucosa (Murphy et al., 1987, J. Infect. Dis. 5:723-731). Colonization with a particular strain may persist for weeks to months, and most individuals remain 25 asymptomatic throughout this period (Spinola et al., 1986, I. Infect. Dis. 154:100-109). However, in certain circumstances colonization will be followed contiguous spread within the respiratory tract, resulting in local disease in the middle ear, the 30 sinuses, the conjunctiva, or the lungs. Alternatively, on occasion bacteria will penetrate the nasopharyngeal epithelial barrier and enter the bloodstream.

In vitro observations and animal studies suggest that bacterial surface appendages called pili (or fimbriae) play an important role in H. influenzae colonization. In 1982 two groups reported a correlation between and increased attachment to human oropharyngeal epithelial cells and erythrocytes (Guerina et al., supra; Pichichero et al., supra). investigators have demonstrated that anti-pilus antibodies block in vitro attachment by piliated H. influenzae (Forney et al., 1992, J. Infect. 165:464-470; van Alphen et al., 1988, Infect. Immun. 56:1800-1806). Recently Weber et al. insertionally inactivated the pilus structural gene influenzae type b strain and thereby eliminated expression of pili; the resulting mutant exhibited a reduced capacity for colonization of year-old monkeys (Weber et al., 1991, Infect. Immun. 59:4724-4728).

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A number of reports suggest that nonpilus factors also facilitate Haemophilus colonization. Using the human nasopharyngeal organ culture model, Farley et al. (1986, J. Infect. Dis. 161:274-280) and Loeb et al. (1988, Infect. Immun. 49:484-489) noted that nonpiliated type b strains were capable of mucosal attachment. Read and coworkers made similar observations upon examining nontypable strains in a model that employs nasal turbinate tissue in organ culture (1991, J. Infect. Dis. 163:549-558). In the monkey colonization study by Weber et al. (1991, supra), nonpiliated organisms retained a capacity for colonization, though at reduced densities; moreover, among monkeys originally infected with the piliated strain, virtually all organisms recovered from the nasopharynx were nonpiliated. All of observations are consistent with the finding that nasopharyngeal isolates from children colonized with H.

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influenzae are frequently nonpiliated (Mason et al., 1985, Infect. Immun. 49:98-103; Brinton et al., 1989, Pediatr. Infect. Dis. J. 8:554-561).

Previous studies have shown that H. influenzae are capable of entering (invading) cultured human epithelial 5 cells via a pili-independent mechanism (St. Geme and Falkow, 1990, supra; St. Geme and Falkow, 1991, supra). Although H. influenzae is not generally considered an intracellular parasite, a recent report suggests that these in vitro findings may have an in vivo correlate 10 (Forsgren et al., 1994, supra). Forsgren and coworkers examined adenoids from 10 children who had their adenoids removed because of longstanding secretory otitis media or adenoidal hypertrophy. In all 10 cases there were viable intracellular H. influenzae. Electron 15 microscopy demonstrated that these organisms were concentrated in the reticular crypt epithelium and in macrophage-like cells in the subepithelial layer of One possibility is that bacterial entry into host cells provides a mechanism for evasion of the local 20 immune response, thereby allowing persistence in the respiratory tract.

Thus, a vaccine for the therapeutic and prophylactic treatment of Haemophilus infection is desirable. Accordingly, it is an object of the present invention to provide for recombinant Haemophilus Adherence and Penetration (HAP) proteins and variants thereof, and to produce useful quantities of these HAP proteins using recombinant DNA techniques.

It is a further object of the invention to provide recombinant nucleic acids encoding HAP proteins, and

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expression vectors and host cells containing the nucleic acid encoding the HAP protein.

An additional object of the invention is to provide monoclonal antibodies for the diagnosis of *Haemophilus* infection.

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A further object of the invention is to provide methods for producing the HAP proteins, and a vaccine comprising the HAP proteins of the present invention. Methods for the therapeutic and prophylactic treatment of Haemophilus infection are also provided.

### SUMMARY OF THE INVENTION

In accordance with the foregoing objects, the present invention provides recombinant HAP proteins, and isolated or recombinant nucleic acids which encode the HAP proteins of the present invention. Also provided are expression vectors which comprise DNA encoding a HAP protein operably linked to transcriptional and translational regulatory DNA, and host cells which contain the expression vectors.

- The invention provides also provides methods for producing HAP proteins which comprises culturing a host cell transformed with an expression vector and causing expression of the nucleic acid encoding the HAP protein to produce a recombinant HAP protein.
- The invention also includes vaccines for Haemophilus influenzae infection comprising an HAP protein for prophylactic or therapeutic use in generating an immune response in a patient. Methods of treating or

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preventing Haemophilus influenzae infection comprise administering a vaccine.

#### BRIEF DESCRIPTION OF THE DRAWINGS

Figures 1A and 1B depict light micrographs of H. influenzae strains DB117(pGJB103) and DB117(pN187) incubated with Chang epithelial cells. Bacteria were incubated with an epithelial monolayer for 30 minutes before rinsing and straining with Giemsa stain. Figure 1A: H. influenzae strain DB117 carrying cloning vector alone (pGJB103); Figure 1B: H. influenzae strain DB117 harboring recombinant plasmid pH187. Bar represents 3.5  $\mu$ m.

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2C and 2D depict thin section Figures 2A, 2B, transmission electron micrographs demonstrating interaction between H. influenzae strains N187 and DB117 (pN187) with Chang epithelial cells. Bacteria were incubated with epithelial monolayers for four hours before rinsing and processing for examination by transmission electron microscopy. Figure 2A: strain N187 associated with the epithelial cell surface and present in an intracellular location; Figure 2B: H. influenzae DB117 (pH187) in intimate contact with the epithelial cell surface; Figure 2C: strain DB117(pN187) in the process of entering an epithelial cell; Figure 2D: strain DB117(pN187) present in an intracellular location. Bar represents 1 µm.

Figure 3 depicts outer membrane protein profiles of various strains. Outer membrane proteins were isolated on the basis of sarcosyl insolubility and resolved on a 10% SDS-polyacrylamide gel. Proteins were visualized by staining with Coomassie blue. Lane 1, H. influenzae

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strain DB117(pGJB103); lane 2, strain DB117(pN187); lane 3, strain DB117(pJS106); lane 4, E. coli HB101(pGJB103); lane 5, HB101(pN187). Note novel proteins at ~160 kD and 45 kD marked by asterisks in lanes 2 and 3.

5 Figure 4 depicts a restriction map of pN187 and derivatives and locations of mini-Tn10 kan insertions. pN187 is a derivative of pGJB103 that contains an 8.5-kb Sau3AI fragment of chromosomal DNA from H. influenzae strain N187. Vector sequences are represented by 10 hatched boxes. Letters above top horizontal line indicate restriction enzyme sites: Bg, BglII; C, ClaI; E, EcoRI; P, PstI. Numbers and lollipops above top horizontal line show positions of mini-Tn10 kan insertions; open lollipops represent insertions that 15 have no effect on adherence and invasion, while closed lollipops indicate insertions that eliminate the capacity of pN187 to promote association with epithelial monolayers. Heavy horizontal line with arrow represents location of hap locus within pN187 and direction of 20 transcription. (+): recombinant plasmids that promote adherence and invasion; (-): recombinant plasmids that fail to promote adherence and invasion.

Figure 5 depicts the identification of plasmid-encoded proteins using the bacteriophage T7 expression system. Bacteria were radiolabeled with [35] methionine, and whole cell lysates were resolved on a 10% SDSpolyacrylamide gel. Proteins were visualized by Lane 1, E. coli XL-1 Blue(pT7-7) autoradiography. uninduced; lane 2, XL-1 Blue(pT7-7) induced with IPTG; lane 3, XL-1 Blue(pJS103) uninduced; lane 4, induced with 5, Blue (pJS103) IPTG; lane XL-1 Blue(pJS104) uninduced; lane 6, XL-1 Blue(pJS104) induced with IPTG. The plasmids pJS103 and pJS104 are

derivatives of pT7-7 that contain the 6.5-kb PstI fragment from pN187 in opposite orientations. Asterisk indicates overexpressed protein in XL-1 Blue(pJS104).

Figures 6A, 6B, and 6C depict the nucleotide sequence and predicted amino acid sequence of hap gene. Putative -10 and -35 sequences 5' to the hap coding sequence are underlined; a putative rho-independent terminator 3' to the hap stop codon is indicated with inverted arrows. The first 25 amino acids of the protein, which are boxed, represent the signal sequence.

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Figures 7A, 7B, 7C, 7D, 7E, 7F, 7G, and 7H depict a sequence comparison of the hap product and the cloned H. influenzae IgAl proteases. Amino acid homologies between the deduced hap gene product and the iga gene products from H. influenzae HK368, HK61, HK393, and HK793 are shown. Dashes indicate gaps introduced in the sequences in order to obtain maximal homology. A consensus sequence for the five proteins is shown on the lower line. The conserved serine-type protease catalytic domain is underlined, and the common active site serine is denoted by an asterisk. The conserved cysteines are also indicated by asterisks.

Figure 8 depicts the IgA1 protease activity assay. Culture supernatants were assayed for the ability to cleave IgA1. Reaction mixtures were resolved on a 10% SDS-polyacrylamide gel and then transferred to a nitrocellulose membrane. The membrane was probed with antibody against human IgA1 heavy chain. Lane 1, H. influenzae strain N187; lane 2, strain DB117(pGJB103); lane 3, strain DB117(pN187). The cleavage product patterns suggest that strain N187 contains a type 2 IgA1 protease while strains DB117(pGJB103) and DB117(pN187)

contain a type 1 enzyme. The upper band of ~70-kD seen with the DB117 derivatives represents intact IgAl heavy chain.

Figures 9A and depict southern analysis 9B chromosomal DNA from strain H. influenzae N187, probing 5 with hap versus iga. DNA fragments were separated on a 0.7% agarose gel and transferred bidirectionally to nitrocellulose membranes prior to probing with either hap or iga. Lane 1, N187 chromosomal DNA digested with 10 EcoRI; lane 2, N187 chromosomal DNA digested with BglII; lane 3, N187 chromosomal DNA digested with BamHI; lane 4, the 4.8-kb ClaI-PstI fragment from pN187 that contains the intact hap gene. Figure 9A: Hybridization with the 4.8-kb ClaI-PstI fragment containing the hap 15 gene; Figure 9B: hybridization with the iga gene from H. influenzae strain Rd, carried as a 4.8-kb ClaI-EcoRI fragment in pVD116.

Figure 10 depicts a SDS-polyacrylamide gel of secreted proteins. Bacteria were grown to late log phase, and 20 culture supernatants were precipitated with trichloroacetic acid and then resolved on a 10% SDSpolyacrylamide gel. Proteins were visualized by staining with Coomassie blue. Lane 1, H. influenzae strain DB117(pGJB103); lane 2, DB117(pN187); lane 3, 25 DB117(pJS106); lane 4, DB117(pJS102); lane DB117(pJS105); lane DB117(Tn10-18); 6, lane 7, DB117(Tn10-4'); lane 8, DB117(Tn10-30); lane DB117(Tn10-16); lane 10, DB117(Tn10-10); DB117(Tn10-8); lane 12, N187. Asterisk indicates 110-kD 30 secreted protein encoded by hap.

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# DETAILED DESCRIPTION OF THE INVENTION

The present invention provides novel Haemophilus Adhesion and Penetration (HAP) proteins. In a preferred embodiment, the HAP proteins are from Haemophilus strains, and in the preferred embodiment, from Haemophilus influenza. However, using the techniques outlined below, HAP proteins from other Haemophilus influenzae strains, or from other bacterial species such as Neisseria spp. or Bordetalla spp. may also be obtained.

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A HAP protein may be identified in several ways. A HAP nucleic acid or HAP protein is initially identified by substantial nucleic acid and/or amino acid sequence homology to the sequences shown in Figure 6. Such homology can be based upon the overall nucleic acid or amino acid sequence.

The HAP proteins of the present invention have limited homology to Haemophilus influenzae and N. gonorrhoeae serine-type IgAl proteases. This homology, shown in Figure 7, is approximately 30-35% at the amino acid level, with several stretches showing 55-60% identity, including amino acids 457-549, 399-466, 572-622, and 233-261. However, the homology between the HAP protein and the IgAl protease is considerably lower than the similarity among the IgAl proteases themselves.

In addition, the full length HAP protein has homology to Tsh, a hemagglutinin expressed by an avian *E. coli* strain (Provence and Curtiss 1994, Infect. Immun. 62:1369-1380). The homology is greatest in the N-terminal half of the proteins, and the overall homology is 30.5% homologous. The full length HAP protein also

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has homology with pertactin, a 69 kD outer membrane protein expressed by B. pertussis, with the middle portion of the proteins showing 39% homology. Finally, HAP has 34 - 52% homology with six regions of HpmA, a calcium-independent hemolysin expressed by Proteus mirabilis (Uphoff and Welch, 1990, J. Bacteriol. 172:1206-1216).

As used herein, a protein is a "HAP protein" if the overall homology of the protein sequence to the amino acid sequence shown in Figure 6 is preferably greater than about 40 - 50%, more preferably greater than about 60% and most preferably greater than 80%. embodiments the homology will be as high as about 90 to 95 or 98%. This homology will be determined using standard techniques known in the art, such as the Best Fit sequence program described by Devereux et al., Nucl. Acid Res. 12:387-395 (1984). The alignment may include the introduction of gaps in the sequences to be aligned. In addition, for sequences which contain either more or fewer amino acids than the protein shown in Figure 6, it is understood that the percentage of homology will be determined based on the number of homologous amino acids in relation to the total number of amino acids. Thus, for example, homology of sequences shorter than that shown in Figure 6, as discussed below, will be determined using the number of amino acids in the shorter sequence.

HAP proteins of the present invention may be shorter than the amino acid sequence shown in Figure 6. As shown in the Examples, the HAP protein may undergo post-translational processing similar to that seen for the serine-type IgAl proteases expressed by Haemophilus influenzae and N. gonorrhoeae. These proteases are

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synthesized as preproteins with three functional domains: the N-terminal signal peptide, the protease, and a C-terminal helper domain. Following movement of these proteins into the periplasmic space, the carboxy terminal ß-domain of the proenzyme is inserted into the outer membrane, possibly forming a pore (Poulsen et al., 1989, Infect. Immun. 57:3097-3105; Pohlner et al., 1987, 325:458-462; Klauser et al., 1992, Nature (London). EMBO J. 11:2327-2335; Klauser et al., 1993, J. Mol. Biol. 234:579-593). Subsequently the amino end of the protein is exported through the outer membrane, and autoproteolytic cleavage occurs to result in secretion of the mature 100 to 106-kD protease. The 45 to 56-kD C-terminal ß-domain remains associated with the outer membrane following the cleavage event. As shown in the Examples, the HAP nucleic acid is associated with expression of a 160 kD outer membrane protein. The secreted gene product is an approximately 110 kD protein, with the simultaneous appearance of a 45 kD outer membrane protein. The 45 kD protein appears to correspond to amino acids from about 960 to about 1394 of Figure 6. Any one of these proteins is considered a HAP protein for the purposes of this invention.

Thus, in a preferred embodiment, included within the defintion of HAP proteins are portions or fragments of the sequence shown in Figure 6. The fragments may be fragments of the entire sequence, the 110 kD sequence, or the 45 kD sequence. Generally, the HAP protein fragments may range in size from about 10 amino acids to about 1900 amino acids, with from about 50 to about 1000 amino acids being preferred, and from about 100 to about 500 amino acids also preferred. Particularly preferred fragments are sequences unique to HAP; these sequences have particular use in cloning HAP proteins

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from other organisms or to generate antibodies specific to HAP proteins. Unique sequences are easily identified by those skilled in the art after examination of the HAP protein sequence and comparison to other proteins; for example, by examination of the sequence alignment shown For instance, as compared to the IgA proteases, unique sequences include, but are not limited to, amino acids 11-14, 16-22, 108-120, 155-164, 257-265, 281-288, 318-336, 345-353, 398-416, 684-693, 712-718, 753-761, 871-913, 935-953, 985-1008, 1023-1034, 1067-1076, 1440-1048, 1585-1592, 1631-1639, 1637-1648, 1735-1743, 1863-1871, 1882-1891, 1929-1941, and 1958-1966 (using the numbering of Figure 7). HAP protein fragments which are included within the definition of a HAP protein include N- or C-terminal truncations and deletions which still allow the protein to biologically active; for example, which still exhibit proteolytic activity in the case of the 110 kD putative protease sequence. In addition, when the HAP protein is to be used to generate antibodies, for example as a vaccine, the HAP protein must share at least one epitope or determinant with either the full length protein, the 110 kD protein or the 45 kD protein, shown in Figure 6. In a preferred embodiment, the epitope is unique to the HAP protein; that is, antibodies generated to a unique epitope exhibit little or no cross-reactivity with other proteins. By "epitope" or "determinant" herein is meant a portion of a protein which will generate and/or bind an antibody. Thus, in most instances, antibodies made to a smaller HAP protein will be able to bind to the full length protein.

In some embodiments, the fragment of the HAP protein used to generate antibodies are small; thus, they may

be used as haptens and coupled to protein carriers to generate antibodies, as is known in the art.

Preferably, the antibodies are generated to a portion of the HAP protein which remains attached to the Haemophilus influenzae organism. For example, the HAP protein can be used to vaccinate a patient to produce antibodies which upon exposure to the Haemophilus influenzae organism (e.g. during a subsequent infection) bind to the organism and allow an immune response. Thus, in one embodiment, the antibodies are generated to the roughly 45 kD fragment of the full length HAP protein. Preferably, the antibodies are generated to the portion of the 45 kD fragment which is exposed at the outer membrane.

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In an alternative embodiment, the antibodies bind to the mature secreted 110 kD fragment. For example, as explained in detail below, the HAP proteins of the present invention may be administered therapeutically to generate neutralizing antibodies to the 110 kD putative protease, to decrease the undesirable effects of the 100 kD fragment.

In the case of the nucleic acid, the overall homology of the nucleic acid sequence is commensurate with amino acid homology but takes into account the degeneracy in the genetic code and codon bias of different organisms. Accordingly, the nucleic acid sequence homology may be either lower or higher than that of the protein sequence. Thus the homology of the nucleic acid sequence of Figure 6 is preferably greater than 40%, more preferably greater than about 60% and most preferably greater than

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80%. In some embodiments the homology will be as high as about 90 to 95 or 98%.

In one embodiment, the nucleic acid homology is determined through hybridization studies. Thus, for example, nucleic acids which hybridize under high stringency to all or part of the nucleic acid sequence shown in Figure 6 are considered HAP protein genes. High stringency conditions include washes with 0.1XSSC at 65°C for 2 hours.

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The HAP proteins and nucleic acids of the present invention are preferably recombinant. As used herein, "nucleic acid" may refer to either DNA or RNA. molecules which contain both deoxy- and ribonucleotides. The nucleic acids include genomic DNA, oligonucleotides including sense and anti-sense nucleic acids. Specifically included within the definition of nucleic acid are anti-sense nucleic acids. sense nucleic acid will hybridize to the corresponding non-coding strand of the nucleic acid sequence shown in Figure 6, but may contain ribonucleotides as well as deoxyribonucleotides. Generally, anti-sense nucleic acids function to prevent expression of mRNA, such that a HAP protein is not made, or made at reduced levels. The nucleic acid may be double stranded, stranded, or contain portions of both double stranded or single stranded sequence. By the term "recombinant nucleic acid" herein is meant nucleic acid, originally formed in vitro by the manipulation of nucleic acid by endonucleases, in a form not normally found in nature. Thus an isolated HAP protein gene, in a linear form, or an expression vector formed in vitro by ligating DNA molecules that are not normally joined, are both considered recombinant for the purposes of this

invention. It is understood that once a recombinant nucleic acid is made and reintroduced into a host cell or organism, it will replicate non-recombinantly, i.e. using the <u>in vivo</u> cellular machinery of the host cell rather than <u>in vitro</u> manipulations; however, such nucleic acids, once produced recombinantly, although subsequently replicated non-recombinantly, are still considered recombinant for the purposes of the invention.

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Similarly, a "recombinant protein" is a protein made 10 recombinant techniques, i.e. through expression of a recombinant nucleic acid as depicted A recombinant protein is distinguished from naturally occurring protein by at least one or more 15 characteristics. For example, the protein may be isolated away from some or all of the proteins and compounds with which it is normally associated in its wild type host, or found in the absence of the host cells themselves. Thus, the protein may be partially or substantially purified. The definition includes the 20 production of a HAP protein from one organism in a different organism or host cell. Alternatively, the protein may be made at a significantly higher concentration than is normally seen, through the use of a inducible promoter or high expression promoter, such 25 that the protein is made at increased concentration levels. Alternatively, the protein may be in a form not normally found in nature, as in the addition of an epitope tag or amino acid substitutions, insertions and 30 deletions.

Also included with the definition of HAP protein are HAP proteins from other organisms, which are cloned and expressed as outlined below.

In the case of anti-sense nucleic acids, an anti-sense nucleic acid is defined as one which will hybridize to all or part of the corresponding non-coding sequence of the sequence shown in Figure 6. Generally, the hybridization conditions used for the determination of anti-sense hybridization will be high stringency conditions, such as 0.1XSSC at 65°C.

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Once the HAP protein nucleic acid is identified, it can be cloned and, if necessary, its constituent parts recombined to form the entire HAP protein nucleic acid. Once isolated from its natural source, e.g., contained within a plasmid or other vector or excised therefrom as a linear nucleic acid segment, the recombinant HAP protein nucleic acid can be further used as a probe to identify and isolate other HAP protein nucleic acids. It can also be used as a "precursor" nucleic acid to make modified or variant HAP protein nucleic acids and proteins.

Using the nucleic acids of the present invention which encode HAP protein, a variety of expression vectors are The expression vectors may be either selfreplicating extrachromosomal vectors or vectors which integrate into a host genome. Generally, these include transcriptional expression vectors translational regulatory nucleic acid operably linked to the nucleic acid encoding the HAP protein. "Operably linked" in this context means that the transcriptional and translational regulatory DNA is positioned relative to the coding sequence of the HAP protein in such a manner that transcription is initiated. Generally, this will mean that the promoter and transcriptional initiation or start sequences are positioned 5' to the HAP protein coding region. The transcriptional and

translational regulatory nucleic acid will generally be appropriate to the host cell used to express the HAP protein; for example, transcriptional and translational regulatory nucleic acid sequences from <u>Bacillus</u> will be used to express the HAP protein in <u>Bacillus</u>. Numerous types of appropriate expression vectors, and suitable regulatory sequences are known in the art for a variety of host cells.

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In general, the transcriptional and translational regulatory sequences may include, but are not limited to, promoter sequences, leader or signal sequences, ribosomal binding sites, transcriptional start and stop sequences, translational start and stop sequences, and enhancer or activator sequences. In a preferred embodiment, the regulatory sequences include a promoter and transcriptional start and stop sequences.

Promoter sequences encode either constitutive or inducible promoters. The promoters may be either naturally occurring promoters or hybrid promoters. Hybrid promoters, which combine elements of more than one promoter, are also known in the art, and are useful in the present invention.

In addition, the expression vector may comprise additional elements. For example, the expression vector may have two replication systems, thus allowing it to be maintained in two organisms, for example in mammalian or insect cells for expression and in a procaryotic host for cloning and amplification. Furthermore, for integrating expression vectors, the expression vector contains at least one sequence homologous to the host cell genome, and preferably two homologous sequences which flank the expression construct. The integrating

vector may be directed to a specific locus in the host cell by selecting the appropriate homologous sequence for inclusion in the vector. Constructs for integrating vectors are well known in the art.

In addition, in a preferred embodiment, the expression vector contains a selectable marker gene to allow the selection of transformed host cells. Selection genes are well known in the art and will vary with the host cell used.

10 The HAP proteins of the present invention are produced by culturing a host cell transformed with an expression vector containing nucleic acid encoding a HAP protein, under the appropriate conditions to induce or cause expression of the HAP protein. The conditions 15 appropriate for HAP protein expression will vary with the choice of the expression vector and the host cell. and will be easily ascertained by one skilled in the art through routine experimentation. For example, the use of constitutive promoters in the expression vector will 20 require optimizing the growth and proliferation of the host cell, while the use of an inducible promoter requires the appropriate growth conditions induction. In addition, in some embodiments, the timing the harvest is important. For example, 25 baculoviral systems used in insect cell expression are lytic viruses, and thus harvest time selection can be crucial for product yield.

Appropriate host cells include yeast, bacteria, archebacteria, fungi, and insect and animal cells, including mammalian cells. Of particular interest are <a href="https://doi.org/10.2016/baccharomyces.cerevisiae">Drosophila melangaster</a> cells, <a href="mailto:Saccharomyces.cerevisiae">Saccharomyces cerevisiae</a> and other yeasts, <a href="mailto:E.coli">E.coli</a>, <a href="Baccharomyces.cerevisiae">Baccharomyces.cerevisiae</a> and <a href="mailto:E.coli">E.coli</a>, <a href="Baccharomyces.cerevisiae">Baccharomyces.cerevisiae</a> and <a href="mailto:E.coli">E.coli</a>, <a href="mailto:Baccharomyces.cerevisiae">Baccharomyces.cerevisiae</a> and <a href="mailto:E.coli">Baccharomyces.cerevisiae</a> and <a href="mailto:E.coli">E.coli</a>, <a href="mailto:Baccharomyces.cerevisiae</a> and <a href="mailto:Baccharomyces.cerevisiae</a> and <a href="mailto:E.coli">E.coli</a>, <a href="mailto:Bacc

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C129 cells, 293 cells, Neurospora, BHK, CHO, COS, and HeLa cells, immortalized mammalian myeloid and lymphoid cell lines.

In a preferred embodiment, HAP proteins are expressed in bacterial systems. Bacterial expression systems are well known in the art.

A suitable bacterial promoter is any nucleic acid sequence capable of binding bacterial RNA polymerase and initiating the downstream (3') transcription of the coding sequence of HAP protein into mRNA. A bacterial promoter has a transcription initiation region which is usually placed proximal to the 5' end of the coding sequence. This transcription initiation typically includes an RNA polymerase binding site and a transcription initiation site. Sequences encoding metabolic pathway enzymes provide particularly useful promoter sequences. Examples include promoter sequences derived from sugar metabolizing enzymes, such as galactose, lactose and maltose, and sequences derived from biosynthetic enzymes such as tryptophan. Promoters from bacteriophage may also be used and are known in the In addition, synthetic promoters and hybrid promoters are also useful; for example, the tac promoter is a hybrid of the trp and lac promoter sequences. Furthermore, a bacterial promoter can include naturally occurring promoters of non-bacterial origin that have the ability to bind bacterial RNA polymerase and initiate transcription.

In addition to a functioning promoter sequence, an efficient ribosome binding site is desirable. In E. coli, the ribosome binding site is called the Shine-Delgarno (SD) sequence and includes an initiation codon

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and a sequence 3-9 nucleotides in length located 3 - 11 nucleotides upstream of the initiation codon.

The expression vector may also include a signal peptide sequence that provides for secretion of the HAP protein in bacteria. The signal sequence typically encodes a signal peptide comprised of hydrophobic amino acids which direct the secretion of the protein from the cell, as is well known in the art. The protein is either secreted into the growth media (gram-positive bacteria) or into the periplasmic space, located between the inner and outer membrane of the cell (gram-negative bacteria).

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The bacterial expression vector may also include a selectable marker gene to allow for the selection of bacterial strains that have been transformed. Suitable selection genes include genes which render the bacteria resistant to drugs such as ampicillin, chloramphenicol, erythromycin, kanamycin, neomycin and tetracycline. Selectable markers also include biosynthetic genes, such as those in the histidine, tryptophan and leucine biosynthetic pathways.

These components are assembled into expression vectors. Expression vectors for bacteria are well known in the art, and include vectors for Bacillus subtilis, E. coli, Streptococcus cremoris, and Streptococcus lividans, among others.

The bacterial expression vectors are transformed into bacterial host cells using techniques well known in the art, such as calcium chloride treatment, electroporation, and others.

In one embodiment, HAP proteins are produced in insect Expression vectors for the transformation of insect cells, and in particular, baculovirus-based expression vectors, are well known in the art. Briefly, baculovirus is a very large DNA virus which produces its coat protein at very high levels. Due to the size of the baculoviral genome, exogenous genes must be placed in the viral genome by recombination. Accordingly, the components of the expression system include: a transfer vector, usually a bacterial plasmid, which contains both a fragment of the baculovirus genome, and a convenient restriction site for insertion of the HAP protein; a wild type baculovirus with a sequence homologous to the baculovirus-specific fragment in the transfer vector (this allows for the homologous recombination of the heterologous gene into the baculovirus genome); and appropriate insect host cells and growth media.

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Mammalian expression systems are also known in the art and are used in one embodiment. A mammalian promoter is any DNA sequence capable of binding mammalian RNA polymerase and initiating the downstream transcription of a coding sequence for HAP protein into mRNA. A promoter will have a transcription initiating region, which is usually place proximal to the 5' end of the coding sequence, and a TATA box, using a located base pairs upstream of the transcription initiation site. The TATA box is thought to direct RNA polymerase II to begin RNA synthesis at the correct A mammalian promoter will also contain an upstream promoter element, typically located within 100 to 200 base pairs upstream of the TATA box. An upstream promoter element determines the rate at which transcription is initiated and can act in either orientation. Of particular use as mammalian promoters

are the promoters from mammalian viral genes, since the viral genes are often highly expressed and have a broad host range. Examples include the SV40 early promoter, mouse mammary tumor virus LTR promoter, adenovirus major late promoter, and herpes simplex virus promoter.

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Typically, transcription termination and polyadenylation sequences recognized by mammalian cells are regulatory regions located 3' to the translation stop codon and thus, together with the promoter elements, flank the coding sequence. The 3' terminus of the mature mRNA is formed by site-specific post-translational cleavage and polyadenylation. Examples of transcription terminator and polyadenlytion signals include those derived form SV40.

The methods of introducing exogenous nucleic acid into mammalian hosts, as well as other hosts, is well known in the art, and will vary with the host cell used. Techniques include dextran-mediated transfection, calcium phosphate precipitation, polybrene mediated transfection, protoplast fusion, electroporation, encapsulation of the polynucleotide(s) in liposomes, and direct microinjection of the DNA into nuclei.

In a preferred embodiment, HAP protein is produced in yeast cells. Yeast expression systems are well known in the art, and include expression vectors for Saccharomyces cerevisiae, Candida albicans and C. maltosa, Hansenula polymorpha, Kluyveromyces fraqilis and K. lactis, Pichia quillerimondii and P. pastoris, Schizosaccharomyces pombe, and Yarrowia lipolytica. Preferred promoter sequences for expression in yeast include the inducible GAL1,10 promoter, the promoters from alcohol dehydrogenase, enolase, glucokinase,

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glucose-6-phosphate isomerase, glyceraldehyde-3phosphate-dehydrogenase, hexokinase, phosphofructokinase, 3-phosphoglycerate mutase, pyruvate kinase, and the acid phosphatase gene. Yeast selectable markers include ADE2, HIS4, LEU2, TRP1, and ALG7, which confers resistance to tunicamycin; the G418 resistance gene, which confers resistance to G418; and the CUP1 gene, which allows yeast to grow in the presence of copper ions.

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10 recombinant HAP protein may be expressed intracellularly or secreted. The HAP protein may also be made as a fusion protein, using techniques well known in the art. Thus, for example, if the desired epitope is small, the HAP protein may be fused to a carrier protein to form an immunogen. Alternatively, the HAP 15 protein may be made as a fusion protein to increase expression.

Also included within the definition of HAP proteins of the present invention are amino acid sequence variants. These variants fall into one or more of three classes: 20 substitutional, insertional or deletional variants. These variants ordinarily are prepared by site specific mutagenesis of nucleotides in the DNA encoding the HAP protein, using cassette mutagenesis or other techniques well known in the art, to produce DNA encoding the variant, and thereafter expressing the recombinant cell culture as outlined above. variant HAP protein fragments having up to about 100-150 residues may be prepared by in vitro synthesis using established techniques. Amino acid sequence variants are characterized by the predetermined nature of the variation, a feature that sets them apart from naturally occurring allelic or interspecies variation of the HAP

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protein amino acid sequence. The variants typically exhibit the same qualitative biological activity as the naturally occurring analogue, although variants can also be selected which have modified characteristics as will be more fully outlined below.

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While the site or region for introducing an amino acid sequence variation is predetermined, the mutation per se need not be predetermined. For example, in order to optimize the performance of a mutation at a given site, random mutagenesis may be conducted at the target codon or region and the expressed HAP protein variants screened for the optimal combination of desired activity. Techniques for making substitution mutations at predetermined sites in DNA having a known sequence are well known, for example, M13 primer mutagenesis. Screening of the mutants is done using assays of HAP protein activities; for example, mutated HAP genes are placed in HAP deletion strains and tested for HAP activity, as disclosed herein. The creation of deletion strains, given a gene sequence, is known in the art. For example, nucleic acid encoding the variants may be expressed in a Haemophilus influenzae strain deficient in the HAP protein, and the adhesion and infectivity of the variant Haemophilus influenzae Alternatively, the variant HAP protein may be expressed and its biological characteristics evaluated, example its proteolytic activity.

Amino acid substitutions are typically of single residues; insertions usually will be on the order of from about 1 to 20 amino acids, although considerably larger insertions may be tolerated. Deletions range from about 1 to 30 residues, although in some cases

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deletions may be much larger, as for example when one of the domains of the HAP protein is deleted.

Substitutions, deletions, insertions or any combination thereof may be used to arrive at a final derivative. Generally these changes are done on a few amino acids to minimize the alteration of the molecule. However, larger changes may be tolerated in certain circumstances.

When small alterations in the characteristics of the HAP protein are desired, substitutions are generally made in accordance with the following chart:

#### Chart I

	Original Residue	Exemplary Substitutions
15	Ala Arg Asn	Ser Lys Gln, His
	Asp Cys Gln	Glu Ser Asn
20	Glu Gly His	Asp Pro Asn, Gln
25	Ile Leu Lys Met	Leu, Val Ile, Val Arg, Gln, Glu
	Phe Ser Thr	Leu, Ile Met, Leu, Tyr Thr
30	Trp Tyr Val	Ser Tyr Trp, Phe Ile, Leu

Substantial changes in function or immunological identity are made by selecting substitutions that are less conservative than those shown in Chart I. For

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be made which example, substitutions may significantly affect: the structure of the polypeptide backbone in the area of the alteration, for example the alpha-helical or beta-sheet structure; the charge or hydrophobicity of the molecule at the target site; or the bulk of the side chain. The substitutions which in general are expected to produce the greatest changes in the polypeptide's properties are those in which (a) a hydrophilic residue, e.g. seryl or threonyl, substituted for (or by) a hydrophobic residue, e.q. leucyl, isoleucyl, phenylalanyl, valyl or alanyl; (b) a cysteine or proline is substituted for (or by) any other residue; (c) a residue having an electropositive side chain, e.g. lysyl, arginyl, or histidyl, substituted for (or by) an electronegative residue, e.g. glutamyl or aspartyl; or (d) a residue having a bulky side chain, e.g. phenylalanine, is substituted for (or by) one not having a side chain, e.g. glycine.

The variants typically exhibit the same qualitative biological activity and will elicit the same immune response as the naturally-occurring analogue, although variants also are selected to modify the characteristics of the polypeptide as needed. Alternatively, variant may be designed such that the biological activity of the HAP protein is altered. For example, the proteolytic activity of the larger 110 kD domain of the HAP protein may be altered, through the substitution of the amino acids of the active site. The putative catalytic domain of this protein is GDSGSPMF, with the first serine corresponding to the active site serine characteristic of serine type proteases. The residues of the active site may be individually or simultaneously altered to decrease or eliminate proteolytic activity. This may be done to decrease the toxicity or side

effects of the vaccine. Similarly, the cleavage site between the 45 kD domain and the 100 kD domain may be altered, for example to eliminate proteolytic processing to form the two domains. Putatively this site is at residue 960.

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In a preferred embodiment, the HAP protein is purified or isolated after expression. HAP proteins may be isolated or purified in a variety of ways known to those skilled in the art depending on what other components are present in the sample. Standard purification 10 methods include electrophoretic, molecular, immunological and chromatographic techniques, including ion exchange, hydrophobic, affinity, and reverse-phase HPLC chromatography, and chromatofocusing. For example, the HAP protein may be purified using a standard anti-15 HAP antibody column. Ultrafiltration and diafiltration techniques, in conjunction with protein concentration, are also useful. For general guidance in suitable purification techniques, see Scopes, R., Protein Purification, Springer-Verlag, NY (1982). The degree 20 of purification necessary will vary depending on the use of the HAP protein. In some instances no purification will be necessary.

Once expressed and purified if necessary, the HAP proteins are useful in a number of applications.

For example, the HAP proteins can be coupled, using standard technology, to affinity chromatography columns. These columns may then be used to purify antibodies from samples obtained from animals or patients exposed to the Haemophilus influenzae organism. The purified antibodies may then be used as outlined below.

Additionally, the HAP proteins are useful to make antibodies to HAP proteins. These antibodies find use in a number of applications. In a preferred embodiment, the antibodies are used to diagnose the presence of an Haemophilus influenzae infection in a sample or patient. This will be done using techniques well known in the art; for example, samples such as blood or tissue samples may be obtained from a patient and tested for reactivity with the antibodies, for example using standard techniques such as ELISA. In a preferred embodiment, monoclonal antibodies are generated to the HAP protein, using techniques well known in the art. As outlined above, the antibodies may be generated to the full length HAP protein, or a portion of the HAP protein.

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Antibodies generated to HAP proteins may also be used in passive immunization treatments, as is known in the art.

Antibodies generated to unique sequences of HAP proteins may also be used to screen expression libraries from other organisms to find, and subsequently clone, HAP nucleic acids from other organisms.

In one embodiment, the antibodies may be directly or indirectly labelled. By "labelled" herein is meant a compound that has at least one element, isotope or chemical compound attached to enable the detection of the compound. In general, labels fall into three classes: a) isotopic labels, which may be radioactive or heavy isotopes; b) immune labels, which may be antibodies or antigens; and c) colored or fluorescent dyes. The labels may be incorporated into the compound at any position. Thus, for example, the HAP protein

antibody may be labelled for detection, or a secondary antibody to the HAP protein antibody may be created and labelled.

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In one embodiment, the antibodies generated to the HAP proteins of the present invention are used to purify or separate HAP proteins or the Haemophilus influenzae organism from a sample. Thus for example, antibodies generated to HAP proteins which will bind to the Haemophilus influenzae organism may be coupled, using standard technology, to affinity chromatography columns. These columns can be used to pull out the Haemophilus environmental organism from or tissue Alternatively, antibodies generated to the soluble 110 kD portion of the full-length portion of the protein shown in Figure 7 may be used to purify the 110 kD protein from samples.

In a preferred embodiment, the HAP proteins of the present invention are used as vaccines for prophylactic or therapeutic treatment of a Haemophilus influenzae infection in a patient. By "vaccine" herein is meant an antigen or compound which elicits an immune response in an animal or patient. The vaccine may be administered prophylactically, for example to a patient never previously exposed to the antigen, such that subsequent infection by the Haemophilus influenzae organism is prevented. Alternatively, the vaccine may be administered therapeutically to a patient previously exposed or infected by the Haemophilus influenzae organism. While infection cannot be prevented, in this case an immune response is generated which allows the patient's immune system to more effectively combat the infection. Thus, for example, there may be a decrease or lessening of the symptoms associated with infection.

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A "patient" for the purposes of the present invention includes both humans and other animals and organisms. Thus the methods are applicable to both human therapy and veterinary applications.

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The administration of the HAP protein as a vaccine is done in a variety of ways. Generally, the HAP proteins can be formulated according to known methods to prepare pharmaceutically useful compositions, whereby therapeutically effective amounts of the HAP protein are combined in admixture with a pharmaceutically acceptable Suitable vehicles and vehicle. their carrier formulation are well in the art. Such known compositions will contain an effective amount of the HAP protein together with a suitable amount of vehicle in pharmaceutically prepare acceptable order compositions for effective administration to the host. The composition may include salts, buffers, carrier proteins such as serum albumin, targeting molecules to localize the HAP protein at the appropriate site or tissue within the organism, and other molecules. The composition may include adjuvants as well.

In one embodiment, the vaccine is administered as a single dose; that is, one dose is adequate to induce a sufficient immune response to prophylactically or therapeutically treat a Haemophilus influenzae infection. In alternate embodiments, the vaccine is administered as several doses over a period of time, as a primary vaccination and "booster" vaccinations.

By "therapeutically effective amounts" herein is meant an amount of the HAP protein which is sufficient to induce an immune response. This amount may be different depending on whether prophylactic or therapeutic treatment is desired. Generally, this ranges from about 0.001 mg to about 1 gm, with a preferred range of about 0.05 to about , and the preferred dose being \_\_\_\_\_.

These amounts may be adjusted if adjuvants are used.

The following examples serve to more fully describe the manner of using the above-described invention, as well as to set forth the best modes contemplated for carrying out various aspects of the invention. It is understood that these examples in no way serve to limit the true scope of this invention, but rather are presented for illustrative purposes.

#### **EXAMPLES**

# Example 1 Cloning of the HAP protein

- Bacterial Strains, plasmids, and phage. H. influenzae strain N187 is a clinical isolate that was originally cultivated from the middle ear fluid of a child with acute otitis media. This strain was classified as nontypable based on the absence of agglutination with typing antisera for H. influenzae types a-f (Burroughs Wellcome) and the failure to hybridize with pU038, a plasmid that contains the entire cap b locus (Kroll and Moxon, 1988, J. Bacteriol. 170:859-864).
- H. influenzae strain DB117 is a recl mutant of Rd, a capsule-deficient serotype d strain that has been in the laboratory for over 40 years (Alexander and Leidy, 1951, J. Exp. Med. 83:345-359); DB117 was obtained from G. Barcak (University of Maryland, Baltimore, MD) (Sellow et al., 1968). DB117 is deficient for in vitro adherence and invasion, as assayed below.

H. influenzae strain 12 is the nontypable strain from which the genes encoding the HMW1 and HMW2 proteins were cloned (Barenkamp and Leininger, 1992, Infect. Immun. 60:1302-1313); HMW1 and HMW2 are the prototypic members of a family of nontypable Haemophilus antigenically-related high-molecular-weight adhesive proteins (St. Geme et al., 1993).

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E. coli HB101, which is nonadherent and noninvasive, has been previously described (Sambrook et al., 1989, Molecular cloning: a laboratory manual, 2nd ed. Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y. ). E. coli DH5α was obtained from Bethesda Research Laboratories. E. coli MC1061 was obtained from H. Kimsey (Tufts University, Boston, MA). E. coli XL-1 Blue and the plasmid pBluescript KS- were obtained from Plasmid pT7-7 and phage mGP1-2 were Stratagene. provided by S. Tabor (Harvard Medical School, Boston, MA) (Tabor and Richardson, 1985, Proc. Natl. Acad. Sci. USA. 82:1074-1078). The E. coli-Haemophilus shuttle vector pGJB103 (Tomb et al., 1989, Rd. J. Bacteriol. 171:3796-3802) and phage  $\lambda$ 1105 (Way et al., 1984, Gene. 32:3 69-379) were provided by G. Barcak (University of Maryland, Baltimore, MD). Plasmid pVD116 harbors the IgAl protease gene from H. influenzae strain Rd (Koomey and Falkow, 1984, Infect. Immun. 43:101-107) and was obtained from M. Koomey (University of Michigan, Ann Arbor, MI).

Growth conditions. H. influenzae strains were grown as described (Anderson et al., 1972, J. Clin. Invest. 51:31-38). They were stored at -80°C in brain heart infusion broth with 25% glycerol. E. coli strains were grown on LB agar or in LB broth. They were stored at -80°C in LB broth with 50% glycerol.

For H. influenzae, tetracycline was used in a concentration of 5  $\mu$ g/ml and kanamycin was used in a concentration of 25  $\mu$ g/ml. For E. coli, antibiotics were used in the following concentrations: tetracycline, 12.5  $\mu$ g/ml; kanamycin, 50  $\mu$ g/ml; ampicillin, 100  $\mu$ g/ml.

Recombinant DNA methods. DNA ligations, restriction endonuclease digestions, and gel electrophoresis were performed according to standard techniques (Sambrook et al., 1989, supra). Plasmids were introduced into E. coli strains by either chemical transformation or electroporation, as described (Sambrook et al, 1989, supra; Dower et al., 1988, Nucleic Acids Res. 16:617-6145). In H. influenzae transformation was performed using the MIV method of Herriott et al. (1970, J. Bacteriol. 101:517-524), and electroporation was carried out using the protocol developed for E. coli (Dower et al., 1988, supra).

Construction of genomic library from H. influenzae 20 strain N187. High-molecular-weight chromosomal DNA was prepared from 3 ml of an overnight broth culture of H. influenzae N187 as previously described (Mekalanos, 1983, Cell. 35:253-263). Following partial digestion with Sau3AI, 8 to 12 kb fragments were eluted into DEAE 25 paper (Schleicher & Schuell, Keene, H.H.) and then ligated to BglII-digested calf intestine phosphatasetreated pGJB103. The ligation mixture was electroporated into H. influenzae DB117, and transformants

30 were selected on media containing tetracycline.

Transposon mutagenesis.

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Mutagenesis of plasmid DNA was performed using the mini-Tn10 kan element described by Way et al. (1984, supra). Initially, the appropriate plasmid was introduced into  $E.\ coli\ MC1061$ . The resulting strain was infected with  $\lambda1105$ , which carries the mini-Tn10 kan transposon. Transductants were grown overnight in the presence of kanamycin and an antibiotic to select for the plasmid, and plasmid DNA was isolated using the alkaline lysis method. In order to recover plasmids containing a transposon insertion, plasmid DNA was electroporated into  $E.\ coli\ DH5\alpha$ , plating on media containing kanamycin and the appropriate second antibiotic.

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In order to establish more precisely the region of pN187 involved in promoting interaction with host cells, initially this plasmid was subjected to restriction endonuclease analysis. Subsequently, several subclones were constructed in the vector pGJB103 reintroduced into H. influenzae strain DB117. resulting strains were then examined for adherence and invasion. As summarized in Figure 4, subclones containing either a 3.9-kb PstI-BglII fragment (pJS105) or the adjoining 4.2-kb BglII fragment (pJS102) failed to confer the capacity to associate with Chang cells. In contrast, a subclone containing an insert that included portions of both of these fragments (pJS106) did promote interaction with epithelial monolayers. Transposon mutagenesis performed on pH187 confirmed that the flanking portions of the insert in this plasmid were not required for the adherent/invasive phenotype. the other hand, a transposon insertion located adjacent to the BglII site in pJS106 eliminated adherence and An insertion between the second EcoRI and PstI sites in this plasmid had a similar effect (Figure 4).

### Examination of plasmid-encoded proteins.

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In order to examine plasmid encoded proteins, relevant DNA was ligated into the bacteriophage T7 expression vector pT7-7, and the resulting construct transformed into E. coli XL-1 Blue. Plasmid pT7-7 contains the T7 phage  $\phi$ 10 promoter and ribosomal binding site upstream of a multiple cloning site (Tabor and Richardson, 1985, supra). The T7 promoter was induced by infection with the recombinant M13 phage mGP1-2 and addition of isopropyl- $\beta$ -D-thiogalactopyranoside (final concentration, 1 mM). Phage mGP1-2 contains the gene encoding T7 RNA polymerase, which activates the  $\phi$ 10 promoter in pT7-7 (Tabor and Richardson, 1985, supra).

Like DB117(pN187), strain DB117 carrying pJS106 15 expressed new outer membrane proteins 160-kD and 45-kD in size (Figure 3, lane 3). In order to examine whether the 6.5-kb insert in pJS106 actually encodes these proteins, this fragment of DNA was ligated into the bacteriophage T7 expression vector pT7-7. The resulting plasmid containing the insert in the same orientation 20 as in pN187 was designated pJS104, and the plasmid with the insert in the opposite orientation was designated pJS103. Both pJS104, and p7S103 were introduced into E. coli XL-1 Blue, producing XL-1 Blue(pJS104) and XL-1 Blue (pJS103), respectively. As a negative control, pT7-25 7 was also transformed into XL-1 Blue. The T7 promoter was induced in these three strains by infection with the recombinant M13 phage mGP1-2 and addition of isopropyl- $\beta$ -D-thiogalactopyranoside (final concentration, 1 mM), 30 induced proteins were detected using [355] methionine. As shown in Figure 5, induction of XL-1 Blue(pJS104) resulted in expression of a 160-kD protein and several smaller proteins which presumably represent degradation products. In contrast, when

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Blue (pJS103) and XL-1 Blue (pT7-7) were induced, there was no expression of these proteins. There was no 45-kD protein induced in any of the three strains. This experiment suggested that the 6.5-kb insert present in pJS106 contains the structural gene for the 160-kD outer membrane protein identified in DB117 (pJS106). On the other hand, this analysis failed to establish the origin of the 45-kD membrane protein expressed by DB117 (pJS106).

### 10 Adherence and invasion assays.

Adherence and invasion assays were performed with Chang epithelial cells [Wong-Kilbourne derivative, clone 1-5c-4 (human conjunctiva)], which were seeded into wells of 24-well tissue culture plates as previously described (St. Geme and Falkow, 1990). Adherence was measured after incubating bacteria with epithelial monolayers for 30 minutes as described (St. Geme et al., 1993). Invasion assays were carried out according to our original protocol and involved incubating bacteria with epithelial cells for four hours followed by treatment with gentamicin for two hours (100  $\mu$ g/ml) (St. Geme and Falkow, 1990).

Nucleotide sequence determination and analysis. Nucleotide sequence was determined using a Sequenase kit and double stranded plasmid template. DNA fragments were subcloned into pBluescript KS and sequenced along both strands by primer walking. DNA sequence analysis was performed using the Genetics Computer Group (GCG) software package from the University of Wisconsin (Devereux et al., 1984). Sequence similarity searches were carried out using the BLAST program of the National Center for Biotechnology Information (Altschul et al., 1990, J. Mol. Biol. 215:403-410). The DNA sequence

described here will be deposited in the EMBL/GenBank/DDBJ Nucleotide Sequence Data Libraries.

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Based on the our subcloning results, we reasoned that the central BgIII site in pH187 was positioned within an open reading frame. Examination of a series of mini-Tn10 kan mutants supported this conclusion (Figure 4). Consequently, we sequenced DHA on either side of this BglII site and identified a 4182 bp gene, which we have designated hap for <u>H</u>aemophilus <u>a</u>dherence and <u>p</u>enetration (Figure 6). This gene encodes a 1394 amino acid polypeptide, which we have called Hap, with a calculated molecular mass of 155.4-kD, in good agreement with the molecular mass of the larger of the two novel outer membrane proteins expressed by DB117(pN187) and the protein expressed after induction of XL-1 Blue/pJS104. The hap gene has a G+C content of 39.1%, similar to the published estimate of 38.7% for the whole genome (Kilian, 1976, J. Gen. Microbiol. 93:9-62). Putative -10 and -35 promoter sequences are present upstream of the initiation codon. A consensus ribosomal binding site is lacking. A sequence similar to a rhoindependent transcription terminator is beginning 39 nucleotides beyond the stop codon and contains interrupted inverted repeats with the potential for forming a hairpin structure containing a loop of three bases and a stem of eight bases. Similar to the situation with typical E. coli terminators, this structure is followed by a stretch rich in T residues. Analysis of the predicted amino acid sequence suggested the presence of a 25 amino acid signal peptide at the amino terminus. This region has characteristics typical of procaryotic signal peptides, with three positive Hterminal charges, a central hydrophobic region, and alanine residues at positions 23 and 25 (-3 and -1

relative to the putative cleavage site) (von Heijne, 1984, J. Mol. Biol. 173:243-251).

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Comparison of the deduced amino acid sequence of Hap with other proteins. A protein sequence similarity search was performed with the predicted amino acid sequence using the BLAST network service of the National Center for Biotechnology Information (Altschul et al., This search revealed homology with the 1990, supra). proteases of Н. influenzae and gonorrhoeae. Alignment of the derived amino acid sequences for the hap gene product and the proteases from four different H. influenzae strains revealed homology across the extent of the proteins (Figure 7), with several stretches showing 55-60% identity and 70-80% similarity. Similar levels of homology were noted between the hap product and the IgAl protease from N. gonorrhoeae strain MS11. homology includes the region identified as the catalytic site of the IgAl proteases, which is comprised of the sequence GDSGSPLF, where 2 is the active site serine characteristic of serine proteases (Brenner, 1988, Nature (London). 334:528-530; Poulsen et al., 1992, J. Bacteriol. 174:2913-2921). In the case of Hap, the corresponding sequence is GDSGSPMF. The hap product also contains two cysteines corresponding to the cysteines proposed to be important in forming the catalytic domain of the IgA proteases (Pohlner et al., 1987, supra). Overall there is 30-35% identity and 51-55% similarity between the hap gene product and the H. influenzae and N. gonorrhoeae IqA proteases.

The deduced amino acid sequence encoded by hap was also found to contain significant homology to Tsh, a hemagglutinin expressed by an avian E. coli strain

(Provence and Curtiss, 1994, supra). This homology extends throughout both proteins but is greatest in the H-terminal half of each. Overall the two proteins are 30.5% identical and 51.6% similar. Tsh is also synthesized as a preprotein and is secreted as a smaller form; like the IgAl proteases and perhaps Hap, a carboxy terminal peptide remains associated with the outer membrane (D. Provence, personal communication). While this protein is presumed to have proteolytic activity, substrate has not yet been determined. Interestingly, Tsh was first identified on the basis of its capacity to promote agglutination of erythrocytes. Thus Hap and Tsh are possibly the first members of a novel class of adhesive proteins that are processed analogously to the IgA1 proteases.

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Homology was also noted with pertactin, a 69-kD outer membrane protein expressed by B. pertussis (Charles et al., 1989, Proc. Natl. Acad. Sci. USA. 86:3554-3558). The middle portions of these two molecules are 39% identical and nearly 60% similar. This protein contains the amino acid triplet arginine-glycine-aspartic acid (RGD) and has been shown to promote attachment to cultured mammalian cells via this sequence (Leininger et al., 1991, Proc. Natl. Acad. Sci. USA. 88:345-349). Although Bordetella species are not generally considered intracellular parasites, work by Ewanowich and coworkers indicates that these respiratory pathogens are capable of in vitro entry into human epithelial cells (Ewanowich et al., 1989, Infect. Immun. 57:2698-2704; Ewanowich et al., 1989, Infect. Immun. 57:1240-1247). Leininger et al. reported that preincubation of epithelial monolayers with an RGD-containing peptide derived from the pertactin sequence specifically inhibited B. pertussis entry (Leininger et al., 1992,

Infect. Immun. 60:2380-2385). In addition, these investigators found that coating of Staphylococcus aureus with purified pertactin resulted in more efficient S. aureus entry; the RGD-containing peptide from pertactin inhibited this pertactin-enhanced entry by 75%. Although the hap product lacks an RGD motif, it is possible that Hap and pertactin serve similar biologic functions for H. influenzae and Bordetella species, respectively.

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Additional analysis revealed significant homology (34 to 52% identity, 42 to 70% similarity) with six regions of HpmA, a calcium-independent hemolysin expressed by Proteus mirabilis (Uphoff and Welch, 1990, supra).

The hap locus is distinct from the H. influenzae IgAl protease gene.

Given the degree of similarity between the hap gene product and H. influenzae IgAl protease, we wondered whether we had isolated the IgA1 protease gene of strain To examine this possibility, we performed IqA1 protease activity assays. Among H. influenzae strains. two enzymatically distinct types of IgAl protease have been found (Mulks et al., 1982, J. Infect. Dis. 146:266-Type 1 enzymes cleave the Pro-Ser peptide bond 274). between residues 231 and 232 in the hinge region of human IgAl heavy chain and generate fragments of roughly 28-kD and 31-kD; type 2 enzymes cleave the Pro-Thr bond between residues 235 and 236 in the hinge region and generate 26.5-kD and 32.5-kD fragments. Previous studies of the parent strain from which DB117 was derived have demonstrated that this strain produces a type 1 IgA1 protease (Koomey and Falkow, 1984, supra). As shown in Figure 8, comparison of the proteolytic activities of strain DB117 and strain N187 suggested

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that N187 produces a type 2 IgAl protease. We reasoned that DB117(pN187) might generate a total of four fragments from IgAl protease, consistent with two distinct cleavage specificities. Examination of DB117(pH187) revealed instead that this transformant produces the same two fragments of the IgAl heavy chain as does DB117, arguing that this strain produces only a type 1 enzyme.

In an effort to obtain additional evidence against the possibility that plasmid pH187 contains the N187 IgAl protease gene, we performed a series of Southern blots. As shown in Figure 9, when genomic DNA from strain N187 was digested with EcoRI, BglII, or BamHI and then probed with the hap gene, one set of hybridizing fragments was detected. Probing of the same DNA with the iga gene from H. influenzae strain Rd resulted in a different set of hybridizing bands. Moreover, the iga gene failed to hybridize with a purified 4.8-kb fragment that contained the intact hap gene.

The recombinant plasmid associated with adherence and invasion encodes a secreted protein.

The striking homology between the hap gene product and the Haemophilus and Neisseria IgAl proteases suggested the possibility that these proteins might be processed in a similar manner. The IgAl proteases are synthesized as preproteins with three functional domains: the N-terminal signal peptide, the protease, and a C-terminal helper domain, which is postulated to form a pore in the outer membrane for secretion of the protease (Poulsen et al., 1989, supra; Pohlner et al., 1987, supra). The C-terminal peptide remains associated with the outer membrane following an autoproteolytic cleavage event that results in release of the mature enzyme.

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Consistent with the possibility that the hap gene product follows a similar fate, we found DB117 (pN187) produced a secreted protein approximately 110-kD in size that was absent from DB117(pGJB103) (Figure 10). This protein was also produced by DB117 (pJS106), but not by DB117 (pJ5102) DB117 (pJS105). Furthermore, the two mutants with transposon insertions within the hap coding region were deficient in this protein. In order to determine the relationship between hap and the secreted protein, this protein was transferred to a PVDF membrane and Nterminal amino acid sequencing was performed. Excessive background on the first cycle precluded identification of the first amino acid residue of the free amino terminus. The sequence of the subsequent seven residues was found to be HTYFGID, which corresponds to amino acids 27 through 33 of the hap product.

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The introduction of hap into laboratory strains of E. coli strains was unable to endow these organisms with the capacity for adherence or invasion. In considering these results, it is noteworthy that the E. coli transformants failed to express either the 160-kD or the 45-kD outer membrane protein. Accordingly, they also failed to express the 110-kD secreted protein. explanation for this lack of expression is unclear. One possibility is that the H. influenzae promoter or ribosomal binding site was poorly recognized in E. coli. Indeed the putative -35 sequence upstream of the hap initiation codon is fairly divergent from the  $\sigma70$ consensus sequence, and the ribosomal binding site is unrecognizable. Alternatively, an accessory gene may be required for proper export of the Hap protein, although the striking homology with the IgA proteases,

which are normally expressed and secreted in E. coli, argues against this hypothesis.

In considering the possibility that the hap gene product promotes adherence and invasion by directly binding to a host cell surface structure, it seems curious that the mature protein is secreted from the organism. However, there are examples of other adherence factors that are also secreted. Filamentous hemagglutinin is a 220-kD protein expressed by B. pertussis that mediates in vitro adherence and facilitates natural colonization (Relman et al., 1989, Proc. Natl. Acad. Sci. U.S.A. 86:2637-2641; Kimura et al., 1990, Infect. Immun. 58:7-16). This protein remains surface-associated to some extent but is also released from the cell. The process of Filamentous hemagglutinin secretion involves accessory protein designated FhaC, which appears to be localized to the outer membrane (Willems et al., 1994, Molec. Microbiol. 11:337-347). Similarly, the Ipa proteins implicated in Shigella invasion are also Secretion of these proteins requires the products of multiple genes within the mxi and spa loci (Allaoui et al., 1993, Molec. Microbiol. 7:59-68; Andrews et al., 1991, Infect. Immun. 59:1997-2005; Venkatsan et al., 1992, J. Bacteriol. 174:1990-2001).

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It is conceivable that secretion is simply a consequence of the mechanism for export of the hap gene product to the surface of the organism. However, it is noteworthy that the secreted protein contains a serine-type protease catalytic domain and shows homology with the P. mirobilis hemolysin. These findings suggest that the mature Hap protein may possess proteolytic activity and raise the possibility that Hap promotes interaction with the host cell at a distance by modifying the host cell

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surface. Alternatively, Hap may modify the bacterial surface in order to facilitate interaction with a host cell receptor. It is possible that hap encodes a molecule with dual functions, serving as both adhesin and protease.

### Analysis of outer membrane and secreted proteins.

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Outer membrane proteins were isolated on the basis of sarcosyl insolubility according to the method of Carlone et al. (1986, J. Clin. Microbiol. 24:330-332). Secreted proteins were isolated by centrifuging bacterial cultures at 16,000 g for 10 minutes, recovering the supernatant, and precipitating with trichloroacetic acid in a final concentration of 10%. SDS-polyacrylamide gel electrophoresis was performed as previously described (Laemmli, 1970, Nature (London). 227:680-685).

To identify proteins that might be involved in the interaction with the host cell surface, outer membrane protein profiles for DB117(pN187) and DB117(pGJB103) were compared. As shown in Figure 3, DB117(pN187) expressed two new outer membrane proteins: a high-molecular-weight protein approximately 160-kD in size and a 45-kD protein. *E. coli* HB101 harboring pN187 failed to express these proteins, suggesting an explanation for the observation that HB101(pN187) is incapable of adherence or invasion.

Previous studies have demonstrated that a family of antigenically-related high-molecular-weight proteins with similarity to filamentous hemagglutinin of Bordetella pertussis mediate attachment by nontypable H. influenzae to cultured epithelial cells (St. Geme et al., 1993). To explore the possibility that the gene encoding the strain H187 member of this family was

cloned, whole cell lysates of N187, DB117(pN187), and DB117(pGJB103) were examined by Western immunoblot. Our control strain for this experiment was H. influenzae strain 12. Using a polyclonal antiserum directed against HMW1 and HMW2, the prototypic proteins in this family, we identified a 140-kD protein in strain H187 (not shown). In contrast, this antiserum failed to react with either DB117(pN187) or DB117(pGJB103) (not shown), indicating that pN187 has no relationship to HMW protein expression.

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Determination of amino terminal sequence. Secreted proteins were precipitated with trichloroacetic acid, separated on a 10% SDS-polyacrylamide electrotransferred to a polyvinylidene difluoride (PVDF) membrane (Matsudaira, 1987, J. Biol. Chem. 262:10035-Following staining with Coomassie Brilliant Blue R-250, the 110-kD protein was cut from the PVDF membrane and submitted to the Protein Chemistry Laboratory at Washington University School of Medicine for amino terminal sequence determination. analysis was performed by automated Edman degradation Applied Biosystems Model using an 470A sequencer.

Examination of IgAl protease activity. In order to assess IgAl protease activity, bacteria were inoculated into broth and grown aerobically overnight. Samples were then centrifuged in a microphage for two minutes, and supernatants were collected. A 10  $\mu$ l volume of supernatant was mixed with 16  $\mu$ l of 0.5  $\mu$ g/ml human IgAl (Calbiochem), and chloramphenicol was added to a final concentration of 2  $\mu$ g/ml. After overnight incubation at 37°C, reaction mixtures were electrophoresed on a 10% SDS-polyacrylamide gel, transferred to a nitrocellulose

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membrane, and probed with goat anti-human IgAl heavy chain conjugated to alkaline phosphatase (Kirkegaard & Perry). The membrane was developed by immersion in phosphatase substrate solution (5-bromo-4-chloro-3-indolylphosphate toluidinium-nitro blue tetrazolium substrate system; Kirkegaard & Perry).

Immunoblot analysis. Immunoblot analysis of bacterial whole cell lysates was carried out as described (St. Geme et al., 1991).

Southern hybridization. Southern blotting was performed using high stringency conditions as previously described (St. Geme and Falkow, 1991).

### Microscopy.

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- i. Light microscopy. Samples of epithelial cells with associated bacteria were stained with Giemsa stain and examined by light microscopy as described (St. Geme and Falkow, 1990).
- ii. Transmission electron microscopy. For transmission electron microscopy, bacteria were incubated with epithelial cell monolayers for four hours and were then 20 rinsed four times with PBS, fixed glutaraldehyde/1% osmium tetroxide in 0.1 M sodium phosphate buffer pH 6.4 for two hours on ice, and stained with 0.25% aqueous uranyl acetate overnight. 25 Samples were then dehydrated in graded ethanol solutions and embedded in polybed. Ultrathin sections  $(0.4 \mu m)$ were examined in a Phillips 201c electron microscope.

As shown in Figure 2, DB117(pN187) incubated with monolayers for four hours demonstrated intimate interaction with the epithelial cell surface and was

occasionally found to be intracellular. In a given thin section, invaded cells generally contained one or two intracellular organisms. Of note, intracellular bacteria were more common in sections prepared with strain N187, an observation consistent with results using the gentamicin assay. In contrast, examination of samples prepared with strain DB117 carrying cloning vector alone (pGJB103) failed to reveal internalized bacteria (not shown).

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Having described the preferred embodiments of the present invention it will appear to those of ordinary skill in the art that various modifications may be made to the disclosed embodiments, and that such modifications are intended to be within the scope of the present invention.

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#### SEQUENCE LISTING

(1) GENERAL INFO	RMATION:
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- (i) APPLICANT: Washington University, et al.
- (ii) TITLE OF INVENTION: Haemophilus Adherence and Penetration Protein
- (iii) NUMBER OF SEQUENCES: 9
- (iv) CORRESPONDENCE ADDRESS:
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  - CITY: San Francisco STATE: California (C)
  - (D)
  - COUNTRY: United States (E)
  - (F) ZIP: 94111-4187
- (v) COMPUTER READABLE FORM:
  - (A) MEDIUM TYPE: Floppy disk
  - (B)
  - COMPUTER: IBM PC compatible OPERATING SYSTEM: PC-DOS/MS-DOS (C)
  - SOFTWARE: PatentIn Release #1.0, Version #1.25 (D)
- (vi) CURRENT APPLICATION DATA:
  - (A) APPLICATION NUMBER: PCT/US95/
  - (B) FILING DATE:
  - (C) CLASSIFICATION:
- (vii) PRIOR APPLICATION DATA:
  - APPLICATION NUMBER: US 08/296,791 FILING DATE: 25 AUG 1994 (A)
  - (B)
  - (C) CLASSIFICATION:
- (viii) ATTORNEY/AGENT INFORMATION:
  - NAME: Trecartin, Richard F. (A)
  - REGISTRATION NUMBER: 31,801 (B)
  - (C) REFERENCE/DOCKET NUMBER: FP-59941/RFT
  - TELECOMMUNICATION INFORMATION: (ix)
    - TELEPHONE: (415) 781-1989 (A)
    - TELEFAX: (415) 398-3249 (B)
    - (C) TELEX: 910 277299
- (2) INFORMATION FOR SEQ ID NO:1:
  - SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 4319 base pairs
    - (B) TYPE: nucleic acid
    - (C) STRANDEDNESS: double
    - (D) TOPOLOGY: both
  - (ix) FEATURE:
    - (A) NAME/KEY: CDS
    - (B) LOCATION: 60..4241
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:1:

TCAATAGTCG TTTAACTAGT ATTTTTTAAT ACGAAAAATT ACTTAATTAA ATAAACATT

59

107

ATG AAA AAA ACT GTA TTT CGT CTT AAT TTT TTA ACC GCT TGC ATT TCA

Met Lys Lys Thr Val Phe Arg Leu Asn Phe Leu Thr Ala Cys Ile Ser 10

Leu	GG(	AT/	A GTA e Val 20		CAA Glr	GCG Ala	TGG Trp	GCT Ala 25	я сту	CAC His	ACT Thi	TAT	TTI r Pho	e Gl	ATT y Ile	155
GAT Asp	TAC	CAZ Glr 35	,.	TAT	CGT Arg	GAT Asp	TTT Phe 40	ATE	GAG Glu	AAT Asn	AAA Lys	GGG Gly	Ly:	TTC Ph	C ACA e Thr	203
GTT Val	GGG Gly 50	GCT Ala	CAA Gln	AAT Asn	ATT Ile	AAG Lys 55	vai	TAT	AAC Asn	AAA Lys	CAA Glm 60	Gly	CAA / Glr	TTA	GTT u Val	251
GGC Gly 65	ACA Thr	TCA Ser	ATG Met	ACA Thr	AAA Lys 70	GCC Ala	CCG Pro	ATG Met	ATT Ile	GAT Asp 75	Phe	TCT Ser	GTA Val	GTG Va	TCA Ser 80	299
		,		85	Aza	Deu	vai	GIU	90	GIn	Tyr	Ile	Val	. Sei 95		347
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145			AAA Lys		150	****	GIU	MIG.	Ala	155	Ile	Asp	Met	Thr	Ser 160	539
			GGC Gly	165	••••	- 71	361	Asp	170	rnr	Lys	Tyr	Pro	Glu 175	Arg	587
GTT Val	CGT Arg	ATC Ile	GGC Gly 180	TCT Ser	GGA Gly	CGG Arg	CAG '	TTT Phe 185	TGG Trp	CGA . Arg	AAT Asn	GAT Asp	CAA Gln 190	GAC Asp	AAA Lys	635
GGC Gly	GAC Asp	CAA Gln 195	GTT Val	GCC ( Ala	GGT (	AIA	TAT ( Tyr 200	CAT His	TAT ( Tyr	CTG :	ACA ( Thr	GCT Ala 205	GGC Gly	AAT Asn	ACA Thr	683
	AAT Asn 210	CAG Gln	CGT (	GGA ( Gly .	ura .	GGT 1 Gly . 215	AAT ( Asn	GGA Gly	TAT : Tyr	Ser	TAT 1 Tyr 220	TTG ( Leu	GGA Gly	GGC Gly	GAT Asp	731
GTT / Val . 225	CGT . Arg	AAA Lys	GCG ( Ala	GIY '	GAA : Glu : 230	TAL (	Gly	CCA ' Pro	Leu	CCG 1 Pro 235	ATT ( Ile	GCA ( Ala	GGC (	TCA Ser	AAG Lys 240	779
GGG (	GAC . Asp .	AGT Ser	GGT :	TCT ( Ser ) 245	CCG / Pro 1	ATG 1	TTT / Phe	TTE	TAT ( Tyr ,	SAT ( Asp	GCT ( Ala	GAA 1 Glu	AAA ( Lys	CAA Gln 255	AAA Lys	827
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GAA Glu	AAT Asn	GGG Gly 275	Phe	CAA Gln	TTG Leu	GTT Val	CGC Arg 280	Lys	TCT Ser	TAT Tyr	TTT	GAT Asp 285	Glu	ATT Ile	TTC Phe	923
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TCA Ser	GGA Gly	ATA Ile	CCA Pro	TCA Ser 325	GAA Glu	ATT Ile	AAA Lys	ATT Ile	ACG Thr 330	TTA Leu	GCA Ala	AAT Asn	ATG Met	AGT Ser 335	Leu	1067
CCT Pro	TTG Leu	AAA Lys	GAG Glu 340	AAG Lys	GAT Asp	AAA Lys	GTT Val	CAT His 345	AAT Asn	CCT Pro	AGA Arg	TAT Tyr	GAC Asp 350	GGA Gly	CCT Pro	1115
AAT Asn	ATT Ile	TAT Tyr 355	TCT Ser	CCA Pro	CGT Arg	TTA Leu	AAC Asn 360	AAT Asn	GGA Gly	GAA Glu	ACG Thr	CTA Leu 365	TAT Tyr	TTT Phe	ATG Met	11€3
GAT Asp	CAA Gln 370	AAA Lys	CAA Gln	GGA Gly	TCA Ser	TTA Leu 375	ATC Ile	TTC Phe	GCA Ala	TCT Ser	GAC Asp 380	ATT Ile	AAC Asn	CAA Gln	GGG Gly	1211
GCG Ala 385	GGT Gly	GGT Gly	CTT Leu	TAT Tyr	TTT Phe 390	GAG Glu	GGT Gly	AAT Asn	TTT Phe	ACA Thr 395	GTA Val	TCT Ser	CCA Pro	AAT Asn	TCT Ser 400	1259
AAC Asn	CAA Gln	ACT Thr	TGG Trp	CAA Gln 405	GGA Gly	GCT Ala	GGC Gly	ATA Ile	CAT His 410	GTA Val	AGT Ser	GAA Glu	AAT Asn	AGC Ser 415	ACC Thr	1307
GTT Val	ACT Thr	TGG Trp	AAA Lys 420	GTA Val	AAT Asn	GGC Gly	GTG Val	GAA Glu 425	CAT His	GAT Asp	CGA Arg	CTT Leu	TCT Ser 430	AAA Lys	ATT Ile	1355
GGT Gly	AAA Lys	GGA Gly 435	ACA Thr	TTG Leu	CAC His	GTT Val	CAA Gln 440	GCC Ala	AAA Lys	GGG Gly	GAA Glu	AAT Asn 445	AAA Lys	GGT Gly	TCG Ser	1403
ATC Ile	AGC Ser 450	GTA Val	GGC Gly	GAT Asp	GGT Gly	AAA Lys 455	GTC Val	ATT Ile	TTG Leu	GAG Glu	CAG Gln 460	CAG Gln	GCA Ala	GAC Asp	GAT Asp	1451
CAA Gln 465	Gly	Asn	Lys	Gln	GCC Ala 470	Phe	AGT Ser	GAA Glu	ATT Ile	GGC Gly 475	TTG Leu	GTT Val	AGC Ser	GGC Gly	AGA Arg 480	1499
GGG Gly	ACT Thr	GTT Val	Gln	TTA . Leu 485	AAC Asn	GAT ( Asp	GAT . Asp	AAA Lys	CAA Gln 490	TTT ( Phe	GAT . Asp	ACC Thr	GAT . Asp	AAA Lys 495	TTT Phe	1547
TAT Tyr	TTC Phe	Gly	TTT Phe	CGT ( Arg	GGT ( Gly	GGT ( Gly	CGC ' <b>Arg</b>	TTA Leu 505	GAT ( Asp	CTT : Leu	AAC ( Asn	GGG Gly	CAT His 510	TCA Ser	TTA Leu	1595
ACC Thr	Phe	AAA Lys 515	CGT A	ATC Ile	CAA /	Asn	ACG ( Thr 520	GAC Asp	GAG ( Glu	GGG (	GCA A	ATG Met 525	ATT (	GTG Val	AAC Asn	1643

CAT His	CAA 1 Asr 530	* T177	A ACT	CAA CGlr	GCC Ala	GCT Ala 535	Asr	GTC Val	ACT L Thi	ATT	ACT Thi	r Gl	AAC Y As	GAA n Gl	AGC u Ser	1691
ATT Ile 545	vai	CTA Leu	CCT Pro	TAA T	GGA Gly 550	/ Asn	AAT Asn	ATI	AAT Asr	AAA Lys 555	: Let	GAT Asj	TAC P Ty	AGA	AAA J Lys 560	1739
GAA Glu	ATT Ile	GCC Ala	TAC Tyr	AAC Asn 565	GLY	TGG Trp	TTT Phe	GGC Gly	GAA Glu 570	Thr	GAT Asp	AAA Lys	AAT S As:	AAA n Lys	CAC S His	1787
بزعرن	Gry	Arg	580	ASN	Leu	ille	Tyr	585	Pro	Thr	Thr	Gli	AS] 590	p Arg	ACT Thr	1835
Dea	Deu	595	. ser	GIY	GIY	Inr	600	Leu	Lys	·Gly	Asp	0 Ile	Thi	r Glr	ACA Thr	1883
AAA Lys	GĞT Gly 610	Lys	CTA Leu	TTT	TTC Phe	AGC Ser 615	GGT Gly	AGA Arg	CCG Pro	ACA Thr	CCG Pro 620	His	GCC Ala	TAC Tyr	AAT Asn	1931
CAT His 625	TTA Leu	AAT Asn	AAA Lys	CGT Arg	TGG Trp 630	ser	GAA Glu	ATG Met	GAA Glu	GGT Gly 635	Ile	CCA Pro	CAA Glr	GGC Gly	GAA Glu 640	1979
	V41		Asp	645	Asp	Trp	116	Asn	Arg 650	Thr	Phe	Lys	Ala	655	Asn	2027
TTC Phe	CAA Gln	ATT Ile	AAA Lys 660	GGC Gly	GGA Gly	AGT Ser	GCG Ala	GTG Val 665	GTT Val	TCT Ser	CGC Arg	AAT Asn	GTT Val 670	TCT Ser	TCA Ser	2075
ATT Ile	GAG Glu	GGA Gly 675	AAT Asn	TGG Trp	ACA Thr	GTC Val	AGC Ser 680	AAT Asn	AAT Asn	GCA Ala	AAT Asn	GCC Ala 685	ACA Thr	TTT Phe	GGT Gly	2123
GTT Val	GTG Val 690	CCA Pro	AAT Asn	CAA Gln	CAA Gln	AAT Asn 695	ACC Thr	ATT Ile	TGC Cys	ACG Thr	CGT Arg 700	TCA Ser	GAT Asp	TGG Trp	ACA Thr	2171
GGA Gly 705	TTA Leu	ACG Thr	ACT Thr	TGT Cys	CAA Gln 710	AAA Lys	GTG Val	GAT Asp	TTA Leu	ACC Thr 715	GAT Asp	ACA Thr	AAA Lys	GTT Val	ATT Ile 720	2219
AAT Asn	TCT Ser	ATA Ile	CCA Pro	AAA Lys 725	ACA Thr	CAA Gln	ATC .	AAT Asn	GGC Gly 730	TCT Ser	ATT Ile	AAT Asn	TTA Leu	ACT Thr 735	GAT Asp	2267
AAT Asn	GCA Ala	ACG Thr	GCG Ala 740	AAT Asn	GTT Val	AAA Lys	GGT Gly	TTA Leu 745	GCA . Ala	AAA Lys	CTT Leu	TAA naĄ	GGC Gly 750	AAT Asn	GTC Val	2315
ACT Thr		ACA Thr 755	AAT Asn	CAC . His	AGC Ser	GIN	Phe	ACA Thr	TTA . Leu	AGC . Ser	AAC Asn	AAT Asn 765	GCC Ala	ACC Thr	CAA Gln	2363
	GGC . Gly . 770	AAT . Asn	ATT Ile	CGA Arg	CTT '	TCC ( Ser 775	GAC A	AAT Asn	TCA Z Ser	ACT ( Thr	GCA . Ala 780	ACG Thr	GTG Val	GAT Asp	AAT Asn	2411

-53-

GCA Ala 785	Asr	TTG Leu	AAC Asn	GGT	AAT Asn 790	Val	CAT His	TTA Lev	ACG Thr	GAT Asp 795	Sex	GCT Ala	CAA a Glr	TTT n Phe	TCT Ser 800	2459
TTA Leu	AAA Lys	AAC Asn	AGC Ser	CAT His 805	Phe	TCG Ser	CAC His	CAA Glr	ATT 1le 810	Gln	GGA Gly	GAC Asp	AAA Lys	GGC Gly 815	ACA Thr	2507
ACA Thr	GTG Val	ACG Thr	TTG Leu 820	Glu	AAT Asn	GCG Ala	ACT Thr	TGG Trp 825	Thr	ATG Met	CCT Pro	AGC Ser	GAT Asp 830	Thi	ACA Thr	2555
TTG Leu	CAG Gln	AAT Asn 835	TTA Leu	ACG Thr	CTA Leu	AAT Asn	AAC Asn 840	Ser	ACG Thr	ATC Ile	ACG Thr	TTA Leu 845	Asn	TCA Ser	GCT Ala	2603
TAT Tyr	TCA Ser 850	Ala	AGC Ser	TCA Ser	AAC Asn	AAT Asn 855	ACG Thr	CCA Pro	CGT Arg	CGC Arg	CGT Arg 860	Ser	TTA Leu	GAG Glu	ACG Thr	2651
GAA Glu 865	ACA Thr	ACG Thr	CCA Pro	ACA Thr	TCG Ser 870	GCA Ala	GAA Glu	CAT His	CGT Arg	TTC Phe 875	AAC Asn	ACA Thr	TTG Leu	ACA Thr	GTA Val 880	2699
AAT Asn	GGT Gly	AAA Lys	TTG Leu	AGT Ser 885	GGG Gly	CAA Gln	GGC Gly	ACA Thr	TTC Phe 890	CAA Gln	TTT Phe	ACT Thr	TCA Ser	TCT Ser 895	TTA Leu	2747
TTT Phe	GGC Gly	TAT Tyr	AAA Lys 900	AGC Ser	GAT Asp	AAA Lys	TTA Leu	AAA Lys 905	TTA Leu	TCC Ser	AAT Asn	GAC Asp	GCT Ala 910	GAG Glu	GGC Gly	2795
GAT Asp	TAC Tyr	ATA Ile 915	TTA Leu	TCT Ser	GTT Val	CGC Arg	AAC Asn 920	ACA Thr	GGC Gly	AAA Lys	GAA Glu	CCC Pro 925	GAA Glu	ACC Thr	CTT Leu	2843
GAG Glu	CAA Gln 930	TTA Leu	ACT Thr	TTG Leu	GTT Val	GAA Glu 935	AGC Ser	AAA Lys	GAT Asp	AAT Asn	CAA Gln 940	CCG Pro	TTA Leu	TCA Ser	GAT Asp	2891
AAG Lys 945	CTC Leu	AAA Lys	TTT Phe	ACT Thr	TTA Leu 950	GAA Glu	AAT Asn	GAC Asp	CAC His	GTT Val 955	GAT Asp	GCA Ala	GGT Gly	GCA Ala	TTA Leu 960	2939
CGT Arg	TAT Tyr	AAA Lys	TTA Leu	GTG Val 965	AAG Lys	AAT Asn	GAT Asp	GGC Gly	GAA Glu 970	TTC Phe	CGC Arg	TTG Leu	CAT His	AAC Asn 975	CCA Pro	2987
ATA Ile	AAA Lys	GAG Glu	CAG Gln 980	GAA Glu	TTG Leu	CAC His	AAT Asn	GAT Asp 985	TTA Leu	GTA . Val	AGA Arg	GCA Ala	GAG Glu 990	CAA Gln	GCA Ala	3035
GAA Glu	CGA Arg	ACA Thr 995	TTA ( Leu	GAA Glu	GCC Ala	Lys	CAA Gln 1000	Val	GAA Glu	CCG . Pro	ACT ( Thr	GCT Ala 1009	Lys	ACA Thr	CAA Gln	3083
ACA Thr	GGT Gly 1010	Glu	CCA / Pro	AAA Lys	GTG Val	CGG ' <b>Arg</b> 1015	Ser	AGA Arg	AGA Arg	GCA ( Ala	GCG A Ala 1020	Arg	GCA Ala	GCG Ala	TTT Phe	3131
CCT Pro 1025	Asp	ACC Thr	CTG ( Leu	Pro	GAT Asp 1030	Gln	AGC ( Ser	CTG Leu	TTA . Leu	AAC ( Asn 1035	Ala	TTA Leu	GAA Glu	GCC . Ala	AAA Lys 1040	3179

CAA Gln	GCT Ala	GAA Glu	Leu	Thr 104	; Ala	GAA Glu	ACA Thr	CAA Glr	Lys 105	Ser	AAG Lys	GCA Ala	AAA Lys	ACA Thr 105	Lys	3227
AAA Lys	GTG Val	CGG Arg	TCA Ser 106	Lys	AGA	GCA Ala	GTG Val	TTT Phe 106	: Ser	GAT Asp	CCC Pro	CTG Leu	CTT Leu 107	Asp	CAA Gln	3275
AGC Ser	CTG Leu	TTC Phe 107	Ala	TTA Leu	GAA Glu	GCC Ala	GCA Ala 108	Leu	GAG Glu	GTT Val	ATT Ile	GAT Asp 108	Ala	CCA Pro	CAG Gln	. 3323
CAA Gln	TCG Ser 109	GIU	AAA Lys	GAT Asp	CGT	CTA Leu 109	Ala	CAA Gln	GAA Glu	GAA Glu	GCG Ala 110		AAA Lys	CAA Gln	CGC Arg	3371
110	GIN 5	ьуs	Asp	Leu	11e 111	Ser O	Arg	Тут	Ser	Asn 111	Ser 5		Leu	Ser	Glu 1120	
TTA Leu	TCT Ser	GCA Ala	ACA Tḥr	GTA Val 112	Asn	AGT Ser	ATG Met	CTT	TCT Ser 113	Val	ĊAA Gln	GAT Asp	GAA Glu	TTA Leu 113	Asp	3467
CGT Arg	CTT Leu	TTT Phe	GTA Val 114	Asp	CAA Gln	GCA Ala	CAA Gln	TCT Ser 114	Ala	GTG Val	TGG Trp	ACA Thr	AAT Asn 115	Ile	GCA Ala	3515
GIN	Asp	115	Arg 5	Arg	Tyr	Asp	Ser 116	Asp 0	Ala	Phe	Arg	GCT Ala 116	Tyr 5	Gln	Gln	3563
GIN	1170	o	Asn	Leu	Arg	Gln 117	Ile 5	Gly	Val	Gln	Lys 118	_	Leu	Ala	Asn	3611
1185	Arg	11e	GIY	АТА	119	Phe	Ser	His	Ser	Arg 119	Ser	GAT Asp	Asn	Thr	Phe 1200	3659
Asp	GIU	Gin	Val	Lys 1209	Asn	His	Ala	Thr	Leu 1210	Thr	Met	ATG Met	Ser	Gly 1215	Phe	3707
мта	GIN	Tyr	1220	Trp	GIA	Asp	Leu	Gln 122	Phe 5	Gly	Val	AAC Asn	Val 1230	Gly	Thr	3755
GGA Gly	ATC Ile	AGT Ser 1235	ATA	AGT Ser	AAA Lys	ATG Met	GCT Ala 1240	Glu	GAA Glu	CAA Gln	AGC Ser	CGA Arg 1245	Lys	ATT Ile	CAT His	3803
CGA Arg	AAA Lys 1250	ATA	ATA Ile	AAT Asn	TAT Tyr	GGC Gly 1255	Val	AAT Asn	GCA Ala	AGT Ser	TAT Tyr 1260	CAG ( Gln	TTC ( Phe	CGT ' Arg	TTA Leu	3851
1265	GIn	ren	GIÀ	Ile	Gln 1270	Pro	Tyr	Phe	Gly	Val 1275	Asn	CGC '	Tyr.	Phe	Ile 1280	3899
GAA Glu	CGT Arg	GAA Glu	TAA Asn	TAT Tyr 1285	GID	TCT Ser	GAG Glu	Glu	GTG Val 1290	Arg	GTG . Val	AAA /	Thr	CCT I	Ser	3947

								-	55-								
CTT Leu	GCA Ala	TTT Phe	AAT Asn 130	Arg	TAT Tyr	AAT Asn	GCT Ala	GGC Gly 130	Ile	CGA Arg	GTT Val	GAT Asp	TAT Tyr 131		TTT Phe	:	3995
ACT Thr	CCG Pro	ACA Thr 131	Asp	AAT Asn	ATC Ile	AGC Ser	GTT Val 132	Lys	CCT Pro	TAT Tyr	TTC Phe	TTC Phe 132	Val	AAT Asn	TAT Tyr	4	043
GTT Val	GAT Asp 1330	Val	TCA Ser	AAC Asn	GCT Ala	AAC Asn 133	Val	CAA Gln	ACC Thr	ACG Thr	GTA Val 134	Asn	CTC Leu	ACG Thr	GTG Val	4	1091
TTG Leu 1349	Gln	CAA Gln	CCA Pro	TTT Phe	GGA Gly 135	Arg	TAT Tyr	TGG Trp	CAA Gln	AAA Lys 135	Glu	GTG Val	GGA Gly	TTA Leu	AAG Lys 1360	4	139
GCA Ala	GAA Glu	ATT Ile	TTA Leu	CAT His 1369	Phe	CAA Gln	ATT Ile	TCC Ser	GCT Ala 1370	Phe	ATC Ile	TCA Ser	AAA Lys	TCT Ser 137	Gln	4	187
GGT Gly	TCA Ser	CAA Gln	CTC Leu 1380	Gly	AAA Lys	CAG Gln	CAA Gln	AAT Asn 138	Val	GGC Gly	GTG Val	aaa Lys	TTG Leu 139	GGC GGC	TAT Tyr	4	235
CGT Arg	TGG Trp	ТААА	AATC	AA C	<u>ATA</u>	TTTI	A TO	GTTI	ATTO	ATA	AACA	AGG	TGGG	TCAG	SAT	4	291
CAGA	TCCC	AC C	TTTT	TATE	T C	TAA	LAT									4	319
(2)		RMAT	EQUE (A) (B)	NCE LEN TYF	CHAR IGTH:		RIST 04 an	rics: nino ld		ls							
	(i	i) M	OLEC	ULE	TYPE	: pr	otei	n									

- (11) MODECOME TIPE: procein
- (xi) SEQUENCE DESCRIPTION: SEQ ID NO:2:

Met Lys Lys Thr Val Phe Arg Leu Asn Phe Leu Thr Ala Cys Ile Ser 1 5 10 15

Leu Gly Ile Val Ser Gln Ala Trp Ala Gly His Thr Tyr Phe Gly Ile 20 25 30

Asp Tyr Gln Tyr Tyr Arg Asp Phe Ala Glu Asn Lys Gly Lys Phe Thr

Val Gly Ala Gln Asn Ile Lys Val Tyr Asn Lys Gln Gly Gln Leu Val 50 60

Gly Thr Ser Met Thr Lys Ala Pro Met Ile Asp Phe Ser Val Val Ser 65 70 75 80

Arg Asn Gly Val Ala Ala Leu Val Glu Asn Gln Tyr Ile Val Ser Val 85 90 95

Ala His Asn Val Gly Tyr Thr Asp Val Asp Phe Gly Ala Glu Gly Asn 100 105

Asn Pro Asp Gln His Arg Phe Thr Tyr Lys Ile Val Lys Arg Asn Asn 115 120 125

T	r i	Lys 130	Ly	's Ai	p As	sn Le	u Hi 13	is Pr	7 T	r Gl	u As	Sp As	p Ty	r Hi	s As	n Pro
							•				13	, ,				r Ser 160
						_				1,	U				17	
									10	5				19	0	p Lys
									J				20	5		n Thr
								-				221	0.			/ Asp
	-										د نه	=				Lys 240
										250	,				255	
									20.	,				270	)	Lys
													.285	)		Phe
												300			Gly	
											315	,		•	Gln	320
										330					Ser 335	
									243					350	Gly	
								500					365		Phe	
							3.3					380			Gln	
											395				Asn	400
										410					Ser 415	
									423					430	Lys	
													445		Gly	
												460			<b>Asp</b>	
Gln 465	Gly	/ A:	sn :	Lys	Gln	Ala 470	Phe	Ser	Glu	Ile	Gly 475	Leu	Val	Ser	Gly	Arg 480

Gly Thr Val Gln Leu Asn Asp Asp Lys Gln Phe Asp Thr Asp Lys Phe 485 490 Tyr Phe Gly Phe Arg Gly Gly Arg Leu Asp Leu Asn Gly His Ser Leu 505 Thr Phe Lys Arg Ile Gln Asn Thr Asp Glu Gly Ala Met Ile Val Asn 520 His Asn Thr Thr Gln Ala Ala Asn Val Thr Ile Thr Gly Asn Glu Ser 535 Ile Val Leu Pro Asn Gly Asn Asn Ile Asn Lys Leu Asp Tyr Arg Lys Glu Ile Ala Tyr Asn Gly Trp Phe Gly Glu Thr Asp Lys Asn Lys His 565 570 Asn Gly Arg Leu Asn Leu Ile Tyr Lys Pro Thr Thr Glu Asp Arg Thr Leu Leu Leu Ser Gly Gly Thr Asn Leu Lys Gly Asp Ile Thr Gln Thr Lys Gly Lys Leu Phe Phe Ser Gly Arg Pro Thr Pro His Ala Tyr Asn His Leu Asn Lys Arg Trp Ser Glu Met Glu Gly Ile Pro Gln Gly Glu 630 lle Val Trp Asp His Asp Trp Ile Asn Arg Thr Phe Lys Ala Glu Asn Phe Gln Ile Lys Gly Gly Ser Ala Val Val Ser Arg Asn Val Ser Ser 665 Ile Glu Gly Asn Trp Thr Val Ser Asn Asn Ala Asn Ala Thr Phe Gly Val Val Pro Asn Gln Gln Asn Thr Ile Cys Thr Arg Ser Asp Trp Thr Gly Leu Thr Thr Cys Gln Lys Val Asp Leu Thr Asp Thr Lys Val Ile Asn Ser Ile Pro Lys Thr Gln Ile Asn Gly Ser Ile Asn Leu Thr Asp Asn Ala Thr Ala Asn Val Lys Gly Leu Ala Lys Leu Asn Gly Asn Val 745 Thr Leu Thr Asn His Ser Gln Phe Thr Leu Ser Asn Asn Ala Thr Gln Ile Gly Asn Ile Arg Leu Ser Asp Asn Ser Thr Ala Thr Val Asp Asn 775 Ala Asn Leu Asn Gly Asn Val His Leu Thr Asp Ser Ala Gln Phe Ser Leu Lys Asn Ser His Phe Ser His Gln Ile Gln Gly Asp Lys Gly Thr Thr Val Thr Leu Glu Asn Ala Thr Trp Thr Met Pro Ser Asp Thr Thr

- Leu Gln Asn Leu Thr Leu Asn Asn Ser Thr Ile Thr Leu Asn Ser Ala 835 840 845
- Tyr Ser Ala Ser Ser Asn Asn Thr Pro Arg Arg Ser Leu Glu Thr 850 860
- Glu Thr Thr Pro Thr Ser Ala Glu His Arg Phe Asn Thr Leu Thr Val 865 870 875 880
- Asn Gly Lys Leu Ser Gly Gln Gly Thr Phe Gln Phe Thr Ser Ser Leu 885 890 895
- Phe Gly Tyr Lys Ser Asp Lys Leu Lys Leu Ser Asn Asp Ala Glu Gly 900 905 910
- Asp Tyr Ile Leu Ser Val Arg Asn Thr Gly Lys Glu Pro Glu Thr Leu 915 920 925
- Glu Gln Leu Thr Leu Val Glu Ser Lys Asp Asn Gln Pro Leu Ser Asp 930 935 940
- Lys Leu Lys Phe Thr Leu Glu Asn Asp His Val Asp Ala Gly Ala Leu 945 955 960
- Arg Tyr Lys Leu Val Lys Asn Asp Gly Glu Phe Arg Leu His Asn Pro 965 970 975
- Ile Lys Glu Glu Leu His Asn Asp Leu Val Arg Ala Glu Gln Ala 980 985 990
- Glu Arg Thr Leu Glu Ala Lys Gln Val Glu Pro Thr Ala Lys Thr Gln
  995 1000 1005
- Thr Gly Glu Pro Lys Val Arg Ser Arg Arg Ala Ala Arg Ala Ala Phe 1010 1015 1020
- Pro Asp Thr Leu Pro Asp Gln Ser Leu Leu Asn Ala Leu Glu Ala Lys 1025 1030 1035 1040
- Gln Ala Glu Leu Thr Ala Glu Thr Gln Lys Ser Lys Ala Lys Thr Lys 1045 1050 1055
- Lys Val Arg Ser Lys Arg Ala Val Phe Ser Asp Pro Leu Leu Asp Gln 1060 1065 1070
- Ser Leu Phe Ala Leu Glu Ala Ala Leu Glu Val Ile Asp Ala Pro Gln 1075 1080 1085
- Gln Ser Glu Lys Asp Arg Leu Ala Gln Glu Glu Ala Glu Lys Gln Arg 1090 1095 1100
- Lys Gln Lys Asp Leu Ile Ser Arg Tyr Ser Asn Ser Ala Leu Ser Glu 1105 1110 1115 1120
- Leu Ser Ala Thr Val Asn Ser Met Leu Ser Val Gln Asp Glu Leu Asp 1125 1130 1135
- Arg Leu Phe Val Asp Gln Ala Gln Ser Ala Val Trp Thr Asn Ile Ala 1140 1145 1150
- Gln Asp Lys Arg Arg Tyr Asp Ser Asp Ala Phe Arg Ala Tyr Gln Gln 1155 1160 1165
- Gln Lys Thr Asn Leu Arg Gln Ile Gly Val Gln Lys Ala Leu Ala Asn 1170 1175 1180

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Gly Arg Ile Gly Ala Val Phe Ser His Ser Arg Ser Asp Asn Thr Phe 1185 1190 1195 1200

Asp Glu Gln Val Lys Asn His Ala Thr Leu Thr Met Met Ser Gly Phe 1205 1210 1215

Ala Gln Tyr Gln Trp Gly Asp Leu Gln Phe Gly Val Asn Val Gly Thr 1220 1225 1230

Gly Ile Ser Ala Ser Lys Met Ala Glu Glu Gln Ser Arg Lys Ile His 1235 1240 1245

Arg Lys Ala Ile Asn Tyr Gly Val Asn Ala Ser Tyr Gln Phe Arg Leu 1250 1260

Gly Gln Leu Gly Ile Gln Pro Tyr Phe Gly Val Asn Arg Tyr Phe Ile 1265 1270 1275 1280

Glu Arg Glu Asn Tyr Gln Ser Glu Glu Val Arg Val Lys Thr Pro Ser 1285 1290 1295

Leu Ala Phe Asn Arg Tyr Asn Ala Gly Ile Arg Val Asp Tyr Thr Phe 1300 1305 1310

Thr Pro Thr Asp Asn Ile Ser Val Lys Pro Tyr Phe Phe Val Asn Tyr 1315 1320 1325

Val Asp Val Ser Asn Ala Asn Val Gln Thr Thr Val Asn Leu Thr Val 1330 1335 1340

Leu Gln Gln Pro Phe Gly Arg Tyr Trp Gln Lys Glu Val Gly Leu Lys 1345 1350 1355 1360

Ala Glu Ile Leu His Phe Gln Ile Ser Ala Phe Ile Ser Lys Ser Gln 1365 1370 1375

Gly Ser Gln Leu Gly Lys Gln Gln Asn Val Gly Val Lys Leu Gly Tyr
1380 1385 1390

Arg Trp

- (2) INFORMATION FOR SEQ ID NO:3:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 1541 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: unknown
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:3:

Met Leu Asn Lys Lys Phe Lys Leu Asn Phe Ile Ala Leu Thr Val Ala 1 5 10 15

Tyr Ala Leu Thr Pro Tyr Thr Glu Ala Ala Leu Val Arg Asp Asp Val 20 25 30

Asp Tyr Gln Ile Phe Arg Asp Phe Ala Glu Asn Lys Gly Lys Phe Ser 35 40 45

Val Gly Ala Thr Asn Val Leu Val Lys Asp Lys Asn Asn Lys Asp Leu 50 60

Gly Thr Ala Leu Pro Asn Gly Ile Pro Met Ile Asp Phe Ser Val Val Asp Val Asp Lys Arg Ile Ala Thr Leu Ile Asn Pro Gln Tyr Val Val ... 90 Gly Val Lys His Val Ser Asn Gly Val Ser Glu Leu His Phe Gly Asn Leu Asn Gly Asn Met Asn Asn Gly Asn Ala Lys Ala His Arg Asp Val Ser Ser Glu Glu Asn Arg Tyr Phe Ser Val Glu Lys Asn Glu Tyr Pro Thr Lys Leu Asn Gly Lys Thr Val Thr Thr Glu Asp Gln Thr Gln Lys Arg Arg Glu Asp Tyr Tyr Met Pro Arg Leu Asp Lys Phe Val Thr Glu Val Ala Pro Ile Glu Ala Ser Thr Ala Ser Ser Asp Ala Gly Thr Tyr Asn Asp Gln Asn Lys Tyr Pro Ala Phe Val Arg Leu Gly Ser Gly Ser Gln Phe Ile Tyr Lys Lys Gly Asp Asn Tyr Ser Leu Ile Leu Asn Asn 210 215 220 His Glu Val Gly Gly Asn Asn Leu Lys Leu Val Gly Asp Ala Tyr Thr Tyr Gly Ile Ala Gly Thr Pro Tyr Lys Val Asn His Glu Asn Asn Gly Leu Ile Gly Phe Gly Asn Ser Lys Glu Glu His Ser Asp Pro Lys Gly Ile Leu Ser Gln Asp Pro Leu Thr Asn Tyr Ala Val Leu Gly Asp Ser Gly Ser Pro Leu Phe Val Tyr Asp Arg Glu Lys Gly Lys Trp Leu Phe Leu Gly Ser Tyr Asp Phe Trp Ala Gly Tyr Asn Lys Lys Ser Trp Gln Glu Trp Asn Ile Tyr Lys Ser Gln Phe Thr Lys Asp Val Leu Asn Lys 330 Asp Ser Ala Gly Ser Leu Ile Gly Ser Lys Thr Asp Tyr Ser Trp Ser 345 Ser Asn Gly Lys Thr Ser Thr Ile Thr Gly Gly Glu Lys Ser Leu Asn Val Asp Leu Ala Asp Gly Lys Asp Lys Pro Asn His Gly Lys Ser Val Thr Phe Glu Gly Ser Gly Thr Leu Thr Leu Asn Asn Asn Ile Asp Gln 395 Gly Ala Gly Gly Leu Phe Phe Glu Gly Asp Tyr Glu Val Lys Gly Thr

Ser Asp Asn Thr Trp Lys Gly Ala Gly Val Ser Val Ala Glu Gly Lys Thr Val Thr Trp Lys Val His Asn Pro Gln Tyr Asp Arg Leu Ala Lys Ile Gly Lys Gly Thr Leu Ile Val Glu Gly Thr Gly Asp Asn Lys Gly Ser Leu Lys Val Gly Asp Gly Thr Val Ile Leu Lys Gln Gln Thr Asn Gly Ser Gly Gln His Ala Phe Ala Ser Val Gly Ile Val Ser Gly 485 Arg Ser Thr Leu Val Leu Asn Asp Asp Lys Gln Val Asp Pro Asn Ser 500 505 Ile Tyr Phe Gly Phe Arg Gly Gly Arg Leu Asp Leu Asn Gly Asn Ser Leu Thr Phe Asp His Ile Arg Asn Ile Asp Asp Gly Ala Arg Leu Val Asn His Asn Met Thr Asn Ala Ser Asn Ile Thr Ile Thr Gly Glu Ser Leu Ile Thr Asp Pro Asn Thr Ile Thr Pro Tyr Asn Ile Asp Ala Pro Asp Glu Asp Asn Pro Tyr Ala Phe Arg Arg Ile Lys Asp Gly Gln Leu Tyr Leu Asn Leu Glu Asn Tyr Thr Tyr Tyr Ala Leu Arg Lys Gly Ala Ser Thr Arg Ser Glu Leu Pro Lys Asn Ser Gly Glu Ser Asn Glu 615 Asn Trp Leu Tyr Met Gly Lys Thr Ser Asp Glu Ala Lys Arg Asn Val 630 Met Asn His Ile Asn Asn Glu Arg Met Asn Gly Phe Asn Gly Tyr Phe 650 Gly Glu Glu Gly Lys Asn Asn Gly Asn Leu Asn Val Thr Phe Lys 665 Gly Lys Ser Glu Gln Asn Arg Phe Leu Leu Thr Gly Gly Thr Asn Leu 680 Asn Gly Asp Leu Thr Val Glu Lys Gly Thr Leu Phe Leu Ser Gly Arg 700 . Pro Thr Pro His Ala Arg Asp Ile Ala Gly Ile Ser Ser Thr Lys Lys Asp Pro His Phe Ala Glu Asn Asn Glu Val Val Glu Asp Asp Trp 725 730 Ile Asn Arg Asn Phe Lys Ala Thr Thr Met Asn Val Thr Gly Asn Ala Ser Leu Tyr Ser Gly Arg Asn Val Ala Asn Ile Thr Ser Asn Ile Thr 760

Ala Ser Asn Lys Ala Gln Val His Ile Gly Tyr Lys Thr Gly Asp Thr 775 Val Cys Val Arg Ser Asp Tyr Thr Gly Tyr Val Thr Cys Thr Thr Asp Lys Leu Ser Asp Lys Ala Leu Asn Ser Phe Asn Pro Thr Asn Leu Arg Gly Asn Val Asn Leu Thr Glu Ser Ala Asn Phe Val Leu Gly Lys Ala 825 Asn Leu Phe Gly Thr Ile Gln Ser Arg Gly Asn Ser Gln Val Arg Leu 840 Thr Glu Asn Ser His Trp His Leu Thr Gly Asn Ser Asp Val His Gln 855 Leu Asp Leu Ala Asn Gly His Ile His Leu Asn Ser Ala Asp Asn Ser 875 Asn Asn Val Thr Lys Tyr Asn Thr Leu Thr Val Asn Ser Leu Ser Gly 890 Asn Gly Ser Phe Tyr Tyr Leu Thr Asp Leu Ser Asn Lys Gln Gly Asp Lys Val Val Val Thr Lys Ser Ala Thr Gly Asn Phe Thr Leu Gln Val 920 Ala Asp Lys Thr Gly Glu Pro Asn His Asn Glu Leu Thr Leu Phe Asp Ala Ser Lys Ala Gln Arg Asp His Leu Asn Val Ser Leu Val Gly Asn 955 Thr Val Asp Leu Gly Ala Trp Lys Tyr Lys Leu Arg Asn Val Asn Gly Arg Tyr Asp Leu Tyr Asn Pro Glu Val Glu Lys Arg Asn Gln Thr Val Asp Thr Thr Asn Ile Thr Thr Pro Asn Asn Ile Gln Ala Asp Val Pro 1000 Ser Val Pro Ser Asn Asn Glu Glu Ile Ala Arg Val Asp Glu Ala Pro 1015 1020 Val Pro Pro Pro Ala Pro Ala Thr Pro Ser Glu Thr Thr Glu Thr Val 1035 Ala Glu Asn Ser Lys Gln Glu Ser Lys Thr Val Glu Lys Asn Glu Gln 1050 Asp Ala Thr Glu Thr Thr Ala Gln Asn Arg Glu Val Ala Lys Glu Ala 1065 Lys Ser Asn Val Lys Ala Asn Thr Gln Thr Asn Glu Val Ala Gln Ser 1080 Gly Ser Glu Thr Lys Glu Thr Gln Thr Thr Glu Thr Lys Glu Thr Ala 1095 Thr Val Glu Lys Glu Glu Lys Ala Lys Val Glu Thr Glu Lys Thr Gln

1115

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Glu Val Pro Lys Val Thr Ser Gln Val Ser Pro Lys Gln Glu Gln Ser 1130

- Glu Thr Val Gln Pro Gln Ala Glu Pro Ala Arg Glu Asn Asp Pro Thr 1140 1145 1150
- Val Asn Ile Lys Glu Pro Gln Ser Gln Thr Asn Thr Thr Ala Asp Thr 1160
- Glu Gln Pro Ala Lys Glu Thr Ser Ser Asn Val Glu Gln Pro Val Thr 1175 1180
- Glu Ser Thr Thr Val Asn Thr Gly Asn Ser Val Val Glu Asn Pro Glu 1190 1195
- Asn Thr Thr Pro Ala Thr Thr Gln Pro Thr Val Asn Ser Glu Ser Ser 1205 1210
- Asn Lys Pro Lys Asn Arg His Arg Arg Ser Val Arg Ser Val Pro His
- Asn Val Glu Pro Ala Thr Thr Ser Ser Asn Asp Arg Ser Thr Val Ala 1240
- Leu Cys Asp Leu Thr Ser Thr Asn Thr Asn Ala Val Leu Ser Asp Ala 1255
- Arg Ala Lys Ala Gln Phe Val Ala Leu Asn Val Gly Lys Ala Val Ser 1265 1270 1275
- Gln His Ile Ser Gln Leu Glu Met Asn Asn Glu Gly Gln Tyr Asn Val 1285 1290
- Trp Val Ser Asn Thr Ser Met Asn Lys Asn Tyr Ser Ser Ser Gln Tyr 1300 1305
- Arg Arg Phe Ser Ser Lys Ser Thr Gln Thr Gln Leu Gly Trp Asp Gln 1315 1320
- Thr Ile Ser Asn Asn Val Gln Leu Gly Gly Val Phe Thr Tyr Val Arg 1330 1335
- Asn Ser Asn Asn Phe Asp Lys Ala Thr Ser Lys Asn Thr Leu Ala Gln 1355
- Val Asn Phe Tyr Ser Lys Tyr Tyr Ala Asp Asn His Trp Tyr Leu Gly 1370
- Ile Asp Leu Gly Tyr Gly Lys Phe Gln Ser Lys Leu Gln Thr Asn His 1385
- Asn Ala Lys Phe Ala Arg His Thr Ala Gln Phe Gly Leu Thr Ala Gly 1400
- Lys Ala Phe Asn Leu Gly Asn Phe Gly Ile Thr Pro Ile Val Gly Val
- Arg Tyr Ser Tyr Leu Ser Asn Ala Asp Phe Ala Leu Asp Gln Ala Arg 1430 1435
- Ile Lys Val Asn Pro Ile Ser Val Lys Thr Ala Phe Ala Gln Val Asp 1445 1450
- Leu Ser Tyr Thr Tyr His Leu Gly Glu Phe Ser Val Thr Pro Ile Leu 1465 1470

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Ser Ala Arg Tyr Asp Ala Asn Gln Gly Ser Gly Lys Ile Asn Val Asn 1480

Gly Tyr Asp Phe Ala Tyr Asn Val Glu Asn Gln Gln Gln Tyr Asn Ala 1495

Gly Leu Lys Leu Lys Tyr His Asn Val Lys Leu Ser Leu Ile Gly Gly 1510

Leu Thr Lys Ala Lys Gln Ala Glu Lys Gln Lys Thr Ala Glu Leu Lys

Leu Ser Phe Ser Phe 1540

# (2) INFORMATION FOR SEQ ID NO:4:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 1545 amino acids (E) TYPE: amino acid

  - (D) TOPOLOGY: unknown

# (xi) SEQUENCE DESCRIPTION: SEQ ID NO:4:

Met Leu Asn Lys Lys Phe Lys Leu Asn Phe Ile Ala Leu Thr Val Ala

Tyr Ala Leu Thr Pro Tyr Thr Glu Ala Ala Leu Val Arg Asp Asp Val

Asp Tyr Gln Ile Phe Arg Asp Phe Ala Glu Asn Lys Gly Lys Phe Ser

Val Gly Ala Thr Asn Val Glu Val Arg Asp Lys Asn Asn Arg Pro Leu

Gly Asn Val Leu Pro Asn Gly Ile Pro Met Ile Asp Phe Ser Val Val

Asp Val Asp Lys Arg Ile Ala Thr Leu Val Asn Pro Gln Tyr Val Val

Gly Val Lys His Val Ser Asn Gly Val Ser Glu Leu His Phe Gly Asn

Leu Asn Gly Asn Met Asn Asn Gly Asn Ala Lys Ala His Arg Asp Val

Ser Ser Glu Glu Asn Arg Tyr Tyr Thr Val Glu Lys Asn Glu Tyr Pro

Thr Lys Leu Asn Gly Lys Ala Val Thr Thr Glu Asp Gln Ala Gln Lys

Arg Arg Glu Asp Tyr Tyr Met Pro Arg Leu Asp Lys Phe Val Thr Glu 170

Val Ala Pro Ile Glu Ala Ser Thr Asp Ser Ser Thr Ala Gly Thr Tyr 185

Asn Asn Lys Asp Lys Tyr Pro Tyr Phe Val Arg Leu Gly Ser Gly Thr

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Gln Phe Ile Tyr Glu Asn Gly Thr Arg Tyr Glu Leu Trp Leu Gly Lys Glu Gly Gln Lys Ser Asp Ala Gly Gly Tyr Asn Leu Lys Leu Val Gly Asn Ala Tyr Thr Tyr Gly Ile Ala Gly Thr Pro Tyr Glu Val Asn His Glu Asn Asp Gly Leu Ile Gly Phe Gly Asn Ser Asn Asn Glu Tyr Ile 265 Asn Pro Lys Glu Ile Leu Ser Lys Lys Pro Leu Thr Asn Tyr Ala Val 280 Leu Gly Asp Ser Gly Ser Pro Leu Phe Val Tyr Asp Arg Glu Lys Gly 295 Lys Trp Leu Phe Leu Gly Ser Tyr Asp Tyr Trp Ala Gly Tyr Asn Lys Lys Ser Trp Gln Glu Trp Asn Ile Tyr Lys Pro Glu Phe Ala Glu Lys Ile Tyr Glu Gln Tyr Ser Ala Gly Ser Leu Ile Gly Ser Lys Thr Asp Tyr Ser Trp Ser Ser Asn Gly Lys Thr Ser Thr Ile Thr Gly Gly Glu Lys Ser Leu Asn Val Asp Leu Ala Asp Gly Lys Asp Lys Pro Asn His Gly Lys Ser Val Thr Phe Glu Gly Ser Gly Thr Leu Thr Leu Asn Asn Asn Ile Asp Gln Gly Ala Gly Gly Leu Phe Phe Glu Gly Asp Tyr Glu Val Lys Gly Thr Ser Asp Asn Thr Thr Trp Lys Gly Ala Gly Val Ser 420 Val Ala Glu Gly Lys Thr Val Thr Trp Lys Val His Asn Pro Gln Tyr Asp Arg Leu Ala Lys Ile Gly Lys Gly Thr Leu Ile Val Glu Gly Thr Gly Asp Asn Lys Gly Ser Leu Lys Val Gly Asp Gly Thr Val Ile Leu Lys Gln Gln Thr Asn Gly Ser Gly Gln His Ala Phe Ala Ser Val Gly 490 Ile Val Ser Gly Arg Ser Thr Leu Val Leu Asn Asp Asp Lys Gln Val Asp Pro Asn Ser Ile Tyr Phe Gly Phe Arg Gly Gly Arg Leu Asp Leu 520 Asn Gly Asn Ser Leu Thr Phe Asp His Ile Arg Asn Ile Asp Glu Gly Ala Arg Leu Val Asn His Ser Thr Ser Lys His Ser Thr Val Thr Ile

Thr Gly Asp Asn Leu Ile Thr Asp Pro Asn Asn Val Ser Ile Tyr Tyr 570 Val Lys Pro Leu Glu Asp Asp Asn Pro Tyr Ala Ile Arg Gln Ile Lys 585 Tyr Gly Tyr Gln Leu Tyr Phe Asn Glu Glu Asn Arg Thr Tyr Tyr Ala Leu Lys Lys Asp Ala Ser Ile Arg Ser Glu Phe Pro Gln Asn Arg Gly Glu Ser Asn Asn Ser Trp Leu Tyr Met Gly Thr Glu Lys Ala Asp Ala Gln Lys Asn Ala Met Asn His Ile Asn Asn Glu Arg Met Asn Gly Phe 650 Asn Gly Tyr Phe Gly Glu Glu Glu Gly Lys Asn Asn Gly Asn Leu Asn Val Thr Phe Lys Gly Lys Ser Glu Gln Asn Arg Phe Leu Leu Thr Gly Gly Thr Asn Leu Asn Gly Asp Leu Asn Val Gln Gln Gly Thr Leu Phe 695 Leu Ser Gly Arg Pro Thr Pro His Ala Arg Asp Ile Ala Gly Ile Ser 710 Ser Thr Lys Lys Asp Ser His Phe Ser Glu Asn Asn Glu Val Val Val Glu Asp Asp Trp Ile Asn Arg Asn Phe Lys Ala Thr Asn Ile Asn Val Thr Asn Asn Ala Thr Leu Tyr Ser Gly Arg Asn Val Glu Ser Ile Thr Ser Asn Ile Thr Ala Ser Asn Asn Ala Lys Val His Ile Gly Tyr Lys Ala Gly Asp Thr Val Cys Val Arg Ser Asp Tyr Thr Gly Tyr Val Thr Cys Thr Thr Asp Lys Leu Ser Asp Lys Ala Leu Asn Ser Phe Asn Pro Thr Asn Leu Arg Gly Asn Val Asn Leu Thr Glu Ser Ala Asn Phe Val 825 Leu Gly Lys Ala Asn Leu Phe Gly Thr Ile Gln Ser Arg Gly Asn Ser Gln Val Arg Leu Thr Glu Asn Ser His Trp His Leu Thr Gly Asn Ser, Asp Val His Gln Leu Asp Leu Ala Asn Gly His Ile His Leu Asn Ser Ala Asp Asn Ser Asn Asn Val Thr Lys Tyr Asn Thr Leu Thr Val Asn Ser Leu Ser Gly Asn Gly Ser Phe Tyr Tyr Leu Thr Asp Leu Ser Asn

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Lys Gln Gly Asp Lys Val Val Val Thr Lys Ser Ala Thr Gly Asn Phe 915 920 925

Thr Leu Gln Val Ala Asp Lys Thr Gly Glu Pro Asn His Asn Glu Leu 930 935 940

Thr Leu Phe Asp Ala Ser Lys Ala Gln Arg Asp His Leu Asn Val Ser 945 950 955 960

Leu Val Gly Asn Thr Val Asp Leu Gly Ala Trp Lys Tyr Lys Leu Arg 965 970 975

Asn Val Asn Gly Arg Tyr Asp Leu Tyr Asn Pro Glu Val Glu Lys Arg 980 985 990

Asn Gln Thr Val Asp Thr Thr Asn Ile Thr Thr Pro Asn Asn Ile Gln 995 1000 1005

Ala Asp Val Pro Ser Val Pro Ser Asn Asn Glu Glu Ile Ala Arg Val 1010 1015 1020

Asp Glu Ala Pro Val Pro Pro Pro Ala Pro Ala Thr Pro Ser Glu Thr 1025 1030 1035 1040

Thr Glu Thr Val Ala Glu Asn Ser Lys Gln Glu Ser Lys Thr Val Glu 1045 1050 1055

Lys Asn Glu Gln Asp Ala Thr Glu Thr Thr Ala Gln Asn Arg Glu Val

Ala Lys Glu Ala Lys Ser Asn Val Lys Ala Asn Thr Gln Thr Asn Glu 1075 1080 1085

Val Ala Gln Ser Gly Ser Glu Thr Lys Glu Thr Gln Thr Thr Glu Thr 1090 1095 1100

Lys Glu Thr Ala Thr Val Glu Lys Glu Glu Lys Ala Lys Val Glu Thr 1105 1110 1115 1120

Glu Lys Thr Gln Glu Val Pro Lys Val Thr Ser Gln Val Ser Pro Lys 1125 1130 1135

Gln Glu Gln Ser Glu Thr Val Gln Pro Gln Ala Glu Pro Ala Arg Glu 1140 1145 1150

Asn Asp Pro Thr Val Asn Ile Lys Glu Pro Gln Ser Gln Thr Asn Thr 1155 1160 1165

Thr Ala Asp Thr Glu Gln Pro Ala Lys Glu Thr Ser Ser Asn Val Glu 1170 1175 1180

Gln Pro Val Thr Glu Ser Thr Thr Val Asn Thr Gly Asn Ser Val Val 1185 1190 1195 1200

Glu Asn Pro Glu Asn Thr Thr Pro Ala Thr Thr Gln Pro Thr Val Asn 1205 1210 1215

Ser Glu Ser Ser Asn Lys Pro Lys Asn Arg His Arg Arg Ser Val Arg 1220 1225 1230

Ser Val Pro His Asn Val Glu Pro Ala Thr Thr Ser Ser Asn Asp Arg 1235 1240 1245

Ser Thr Val Ala Leu Cys Asp Leu Thr Ser Thr Asn Thr Asn Ala Val 1250 1255 1260

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Leu Ser Asp Ala Arg Ala Lys Ala Gln Phe Val Ala Leu Asn Val Gly 1265 1270 1275 1280

Lys Ala Val Ser Gln His Ile Ser Gln Leu Glu Met Asn Asn Glu Gly 1285 1290 1295

Gln Tyr Asn Val Trp Val Ser Asn Thr Ser Met Asn Lys Asn Tyr Ser

Ser Ser Gln Tyr Arg Arg Phe Ser Ser Lys Ser Thr Gln Thr Gln Leu 1315 1320 1325

Gly Trp Asp Gln Thr Ile Ser Asn Asn Val Gln Leu Gly Gly Val Phe 1330 1340

Thr Tyr Val Arg Asn Ser Asn Asn Phe Asp Lys Ala Thr Ser Lys Asn 1345 1350 1355 1360

Thr Leu Ala Gln Val Asn Phe Tyr Ser Lys Tyr Tyr Ala Asp Asn His 1365 1376 1375

Trp Tyr Leu Gly Ile Asp Leu Gly Tyr Gly Lys Phe Gln Ser Lys Leu 1380 1385 1390

Gln Thr Asn His Asn Ala Lys Phe Ala Arg His Thr Ala Gln Phe Gly 1395 1400 1405

Leu Thr Ala Gly Lys Ala Phe Asn Leu Gly Asn Phe Gly Ile Thr Pro 1410 1420

Ile Val Gly Val Arg Tyr Ser Tyr Leu Ser Asn Ala Asp Phe Ala Leu 1425 1430 1435 1440

Asp Gln Ala Arg Ile Lys Val Asn Pro Ile Ser Val Lys Thr Ala Phe 1445 1450 1455

Ala Gln Val Asp Leu Ser Tyr Thr Tyr His Leu Gly Glu Phe Ser Val 1460 1465 1470

Thr Pro Ile Leu Ser Ala Arg Tyr Asp Ala Asn Gln Gly Ser Gly Lys 1475 1480 1485

Ile Asn Val Asn Gly Tyr Asp Phe Ala Tyr Asn Val Glu Asn Gln Gln 1490 1500

Gln Tyr Asn Ala Gly Leu Lys Leu Lys Tyr His Asn Val Lys Leu Ser 1505 1510 1515 1520

Leu Ile Gly Gly Leu Thr Lys Ala Lys Gln Ala Glu Lys Gln Lys Thr 1525 1530 1535

Ala Glu Leu Lys Leu Ser Phe Ser Phe 1540 1545

## (2) INFORMATION FOR SEQ ID NO:5:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 1702 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: unknown

## (xi) SEQUENCE DESCRIPTION: SEQ ID NO:5:

Met Leu Asn Lys Lys Phe Lys Leu Asn Phe Ile Ala Leu Thr Val Ala 1 10 15

Tyr Ala Leu Thr Pro Tyr Thr Glu Ala Ala Leu Val Arg Asp Asp Val Asp Tyr Gln Ile Phe Arg Asp Phe Ala Glu Asn Lys Gly Arg Phe Ser Val Gly Ala Thr Asn Val Glu Val Arg Asp Lys Asn Asn His Ser Leu Gly Asn Val Leu Pro Asn Gly Ile Pro Met Ile Asp Phe Ser Val Val Asp Val Asp Lys Arg Ile Ala Thr Leu Ile Asn Pro Gln Tyr Val Val Gly Val Lys His Val Ser Asn Gly Val Ser Glu Leu His Phe Gly Asn Leu Asn Gly Asn Met Asn Asn Gly Asn Asp Lys Ser His Arg Asp Val Ser Ser Glu Glu Asn Arg Tyr Phe Ser Val Glu Lys Asn Glu Tyr Pro Thr Lys Leu Asn Gly Lys Ala Val Thr Thr Glu Asp Gln Thr Gln Lys Arg Arg Glu Asp Tyr Tyr Met Pro Arg Leu Asp Lys Phe Val Thr Glu Val Ala Pro Ile Glu Ala Ser Thr Ala Ser Ser Asp Ala Gly Thr Tyr 185 Asn Asp Gln Asn Lys Tyr Pro Ala Phe Val Arg Leu Gly Ser Gly Thr 200 Gln Phe Ile Tyr Lys Lys Gly Asp Asn Tyr Ser Leu Ile Leu Asn Asn His Glu Val Gly Gly Asn Asn Leu Lys Leu Val Gly Asp Ala Tyr Thr Tyr Gly Ile Ala Gly Thr Pro Tyr Lys Val Asn His Glu Asn Asn Gly Leu Ile Gly Phe Gly Asn Ser Lys Glu Glu His Ser Asp Pro Lys Gly Ile Leu Ser Gln Asp Pro Leu Thr Asn Tyr Ala Val Leu Gly Asp Ser Gly Ser Pro Leu Phe Val Tyr Asp Arg Glu Lys Gly Lys Trp Leu Phe 290 295 300 Leu Gly Ser Tyr Asp Phe Trp Ala Gly Tyr Asn Lys Lys Ser Trp Gln Glu Trp Asn Ile Tyr Lys Pro Glu Phe Ala Lys Thr Val Leu Asp Lys 330 Asp Thr Ala Gly Ser Leu Ile Gly Ser Asn Thr Gln Tyr Asn Trp Asn 345 Pro Thr Gly Lys Thr Ser Val Ile Ser Asn Gly Ser Glu Ser Leu Asn 360

Val Asp Leu Phe Asp Ser Ser Gln Asp Thr Asp Ser Lys Asn Asn 375 His Gly Lys Ser Val Thr Leu Arg Gly Ser Gly Thr Leu Thr Leu Asn Asn Asn Ile Asp Gln Gly Ala Gly Gly Leu Phe Phe Glu Gly Asp Tyr Glu Val Lys Gly Thr Ser Asp Ser Thr Thr Trp Lys Gly Ala Gly Val Ser Val Ala Asp Gly Lys Thr Val Thr Trp Lys Val His Asn Pro Lys Ser Asp Arg Leu Ala Lys Ile Gly Lys Gly Thr Leu Ile Val Glu Gly 455 Lys Gly Glu Asn Lys Gly Ser Leu Lys Val Gly Asp Gly Thr Val Ile 470 Leu Lys Gln Gln Ala Asp Ala Asn Asn Lys Val Lys Ala Phe Ser Gln Val Gly Ile Val Ser Gly Arg Ser Thr Val Val Leu Asn Asp Asp Lys Gln Val Asp Pro Asn Ser Ile Tyr Phe Gly Phe Arg Gly Gly Arg Leu 515 520 525 Asp Ala Asn Gly Asn Asn Leu Thr Phe Glu His Ile Arg Asn Ile Asp Asp Gly Ala Arg Leu Val Asn His Asn Thr Ser Lys Thr Ser Thr Val Thr Ile Thr Gly Glu Ser Leu Ile Thr Asp Pro Asn Thr Ile Thr Pro 570 Tyr Asn Ile Asp Ala Pro Asp Glu Asp Asn Pro Tyr Ala Phe Arg Arg 580 585 590 Ile Lys Asp Gly Gly Gln Leu Tyr Leu Asn Leu Glu Asn Tyr Thr Tyr 600 Tyr Ala Leu Arg Lys Gly Ala Ser Thr Arg Ser Glu Leu Pro Lys Asn Ser Gly Glu Ser Asn Glu Asn Trp Leu Tyr Met Gly Lys Thr Ser Asp Ala Ala Lys Arg Asn Val Met Asn His Ile Asn Asn Glu Arg Met Asn 650 Gly Phe Asn Gly Tyr Phe Gly Glu Glu Glu Gly Lys Asn Asn Gly Asn Leu Asn Val Thr Phe Lys Gly Lys Ser Glu Gln Asn Arg Phe Leu Leu Thr Gly Gly Thr Asn Leu Asn Gly Asp Leu Lys Val Glu Lys Gly Thr Leu Phe Leu Ser Gly Arg Pro Thr Pro His Ala Arg Asp Ile Ala Gly 705 710 715 720

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Ile Ser Ser Thr Lys Lys Asp Gln His Phe Ala Glu Asn Asn Glu Val Val Val Glu Asp Asp Trp Ile Asn Arg Asn Phe Lys Ala Thr Asn Ile Asn Val Thr Asn Asn Ala Thr Leu Tyr Ser Gly Arg Asn Val Ala Asn Ile Thr Ser Asn Ile Thr Ala Ser Asp Asn Ala Lys Val His Ile Gly Tyr Lys Ala Gly Asp Thr Val Cys Val Arg Ser Asp Tyr Thr Gly Tyr Val Thr Cys Thr Thr Asp Lys Leu Ser Asp Lys Ala Leu Asn Ser Phe 810 Asn Ala Thr Asn Val Ser Gly Asn Val Asn Leu Ser Gly Asn Ala Asn Phe Val Leu Gly Lys Ala Asn Leu Phe Gly Thr Ile Ser Gly Thr Gly 840 Asn Ser Gln Val Arg Leu Thr Glu Asn Ser His Trp His Leu Thr Gly 855 Asp Ser Asn Val Asn Gln Leu Asn Leu Asp Lys Gly His Ile His Leu 870 Asn Ala Gln Asn Asp Ala Asn Lys Val Thr Thr Tyr Asn Thr Leu Thr Val Asn Ser Leu Ser Gly Asn Gly Ser Phe Tyr Tyr Leu Thr Asp Leu Ser Asn Lys Gln Gly Asp Lys Val Val Thr Lys Ser Ala Thr Gly Asn Phe Thr Leu Gln Val Ala Asp Lys Thr Gly Glu Pro Thr Lys Asn Glu Leu Thr Leu Phe Asp Ala Ser Asn Ala Thr Arg Asn Asn Leu Asn Val Ser Leu Val Gly Asn Thr Val Asp Leu Gly Ala Trp Lys Tyr Lys Leu Arg Asn Val Asn Gly Arg Tyr Asp Leu Tyr Asn Pro Glu Val Glu 985 Lys Arg Asn Gln Thr Val Asp Thr Thr Asn Ile Thr Thr Pro Asn Asn Ile Gln Ala Asp Val Pro Ser Val Pro Ser Asn Asn Glu Glu Ile Ala 1015 1020 Arg Val Glu Thr Pro Val Pro Pro Pro Ala Pro Ala Thr Pro Ser Glu 1030 1035 Thr Thr Glu Thr Val Ala Glu Asn Ser Lys Gln Glu Ser Lys Thr Val 1045 1050 Glu Lys Asn Glu Gln Asp Ala Thr Glu Thr Thr Ala Gln Asn Gly Glu 1065

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- Val Ala Glu Glu Ala Lys Pro Ser Val Lys Ala Asn Thr Gln Thr Asn 1075 1080 1085
- Glu Val Ala Gln Ser Gly Ser Glu Thr Glu Glu Thr Gln Thr Thr Glu 1090 1095 1100
- Ile Lys Glu Thr Ala Lys Val Glu Lys Glu Glu Lys Ala Lys Val Glu 1105 1110 1115 1120
- Lys Glu Glu Lys Ala Lys Val Glu Lys Asp Glu Ile Gln Glu Ala Pro 1125 1130 1135
- Gln Met Ala Ser Glu Thr Ser Pro Lys Gln Ala Lys Pro Ala Pro Lys 1140 1145 1150
- Glu Val Ser Thr Asp Thr Lys Val Glu Glu Thr Gln Val Gln Ala Gln 1155 1160 1165
- Pro Gln Thr Gln Ser Thr Thr Val Ala Ala Ala Glu Ala Thr Ser Pro 1170 1180
- Asn Ser Lys Pro Ala Glu Glu Thr Gln Pro Ser Glu Lys Thr Asn Ala 1185 1190 1195 1200
- Glu Pro Val Thr Pro Val Val Ser Lys Asn Gln Thr Glu Asn Thr Thr 1205 1210 1215
- Asp Gln Pro Thr Glu Arg Glu Lys Thr Ala Lys Val Glu Thr Glu Lys 1220 1225 1230
- Thr Gln Glu Pro Pro Gln Val Ala Ser Gln Ala Ser Pro Lys Gln Glu 1235 1240 1245
- Gln Ser Glu Thr Val Gln Pro Gln Ala Val Leu Glu Ser Glu Asn Val 1250 1260
- Pro Thr Val Asn Asn Ala Glu Glu Val Gln Ala Gln Leu Gln Thr Gln 1265 1270 1275 1280
- Thr Ser Ala Thr Val Ser Thr Lys Gln Pro Ala Pro Glu Asn Ser Ile 1285 1290 1295
- Asn Thr Gly Ser Ala Thr Ala Ile Thr Glu Thr Ala Glu Lys Ser Asp 1300 1305 1310
- Lys Pro Gln Thr Glu Thr Ala Ala Ser Thr Glu Asp Ala Ser Gln His 1315 1320 1325
- Lys Ala Asn Thr Val Ala Asp Asn Ser Val Ala Asn Asn Ser Glu Ser 1330 1335 1340
- Ser Glu Pro Lys Ser Arg Arg Arg Ser Ile Ser Gln Pro Gln Glu 1345 1350 1355 1360
- Thr Ser Ala Glu Glu Thr Thr Ala Ala Ser Thr Asp Glu Thr Thr Ile 1365 1370 1375
- Ala Asp Asn Ser Lys Arg Ser Lys Pro Asn Arg Arg Ser Arg Arg Ser 1380 1385 1390
- Val Arg Ser Glu Pro Thr Val Thr Asn Gly Ser Asp Arg Ser Thr Val
- Ala Leu Arg Asp Leu Thr Ser Thr Asn Thr Asn Ala Val Ile Ser Asp 1410 1420

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Ala Met Ala Lys Ala Gln Phe Val Ala Leu Asn Val Gly Lys Ala Val 1425 1430 1435 1440

- Ser Gln His Ile Ser Gln Leu Glu Met Asn Asn Glu Gly Gln Tyr Asn 1445 1450 1455
- Val Trp Val Ser Asn Thr Ser Met Asn Glu Asn Tyr Ser Ser Gln 1460 1465 1470
- Tyr Arg Arg Phe Ser Ser Lys Ser Thr Gln Thr Gln Leu Gly Trp Asp 1475 1480 1485
- Gln Thr Ile Ser Asn Asn Val Gln Leu Gly Gly Val Phe Thr Tyr Val 1490 1495 1500
- Arg Asn Ser Asn Asn Phe Asp Lys Ala Ser Ser Lys Asn Thr Leu Ala 1505 1510 1515 1520
- Gln Val Asn Phe Tyr Ser Lys Tyr Tyr Ala Asp Asn His Trp Tyr Leu 1525 1530 1525
- Gly Ile Asp Leu Gly Tyr Gly Lys Phe Gln Ser Asn Leu Lys Thr Asn 1540 1550
- His Asn Ala Lys Phe Ala Arg His Thr Ala Gln Phe Gly Leu Thr Ala 1555 1560 1565
- Gly Lys Ala Phe Asn Leu Gly Asn Phe Gly Ile Thr Pro Ile Val Gly 1570 1575 1580
- Val Arg Tyr Ser Tyr Leu Ser Asn Ala Asn Phe Ala Leu Ala Lys Asp 1585 1590 1595 1600
- Arg Ile Lys Val Asn Pro Ile Ser Val Lys Thr Ala Phe Ala Gln Val 1605 1610 1615
- Asp Leu Ser Tyr Thr Tyr His Leu Gly Glu Phe Ser Val Thr Pro Ile 1620 1630
- Leu Ser Ala Arg Tyr Asp Thr Asn Gln Gly Ser Gly Lys Ile Asn Val 1635 1640 1645
- Asn Gln Tyr Asp Phe Ala Tyr Asn Val Glu Asn Gln Gln Gln Tyr Asn 1650 1660
- Ala Gly Leu Lys Leu Lys Tyr His Asn Val Lys Leu Ser Leu Ile Gly 1665 1670 1675 1680
- Gly Leu Thr Lys Ala Lys Gln Ala Glu Lys Gln Lys Thr Ala Glu Leu 1685 1690 1695

Lys Leu Ser Phe Ser Phe 1700

#### (2) INFORMATION FOR SEQ ID NO:6:

- (i) SEQUENCE CHARACTERISTICS:
  - (A) LENGTH: 1848 amino acids
  - (B) TYPE: amino acid
  - (D) TOPOLOGY: unknown

#### (xi) SEQUENCE DESCRIPTION: SEQ ID NO:6:

Met Leu Asn Lys Lys Phe Lys Leu Asn Phe Ile Ala Leu Thr Val Ala 1 5 10 15

Tyr Ala Leu Thr Pro Tyr Thr Glu Ala Ala Leu Val Arg Asp Asp Val Asp Tyr Gln Ile Phe Arg Asp Phe Ala Glu Asn Lys Gly Lys Phe Ser Val Gly Ala Thr Asn Val Glu Val Arg Asp Lys Lys Asn Gln Ser Leu Gly Ser Ala Leu Pro Asn Gly Ile Pro Met Ile Asp Phe Ser Val Val Asp Val Asp Lys Arg Ile Ala Thr Leu Val Asn Pro Gln Tyr Val Val Gly Val Lys His Val Ser Asn Gly Val Ser Glu Leu His Phe Gly Asn Leu Asn Gly Asn Met Asn Asn Gly Asn Ala Lys Ser His Arg Asp Val Ser Ser Glu Glu Asn Arg Tyr Tyr Thr Val Glu Lys Asn Asn Phe Pro Thr Glu Asn Val Thr Ser Phe Thr Lys Glu Glu Gln Asp Ala Gln Lys Arg Arg Glu Asp Tyr Tyr Met Pro Arg Leu Asp Lys Phe Val Thr Glu 170 Val Ala Pro Ile Glu Ala Ser Thr Ala Asn Asn Asn Lys Gly Glu Tyr Asn Asn Ser Asp Lys Tyr Pro Ala Phe Val Arg Leu Gly Ser Gly Thr 200 Gln Phe Ile Tyr Lys Lys Gly Ser Arg Tyr Gln Leu Ile Leu Thr Glu 210 215 220 Lys Asp Lys Gln Gly Asn Leu Leu Arg Asn Trp Asp Val Gly Gly Asp Asn Leu Glu Leu Val Gly Asn Ala Tyr Thr Tyr Gly Ile Ala Gly Thr Pro Tyr Lys Val Asn His Glu Asn Asn Gly Leu Ile Gly Phe Gly Asn 265 Ser Lys Glu Glu His Ser Asp Pro Lys Gly Ile Leu Ser Gln Asp Pro 280 Leu Thr Asn Tyr Ala Val Leu Gly Asp Ser Gly Ser Pro Leu Phe Val Tyr Asp Arg Glu Lys Gly Lys Trp Leu Phe Leu Gly Ser Tyr Asp Phe Trp Ala Gly Tyr Asn Lys Lys Ser Trp Gln Glu Trp Asn Ile Tyr Lys His Glu Phe Ala Glu Lys Ile Tyr Gln Gln Tyr Ser Ala Gly Ser Leu Ile Gly Ser Asn Thr Gln Tyr Thr Trp Gln Ala Thr Gly Ser Thr Ser 360

Thr Ile Thr Gly Gly Gly Glu Pro Leu Ser Val Asp Leu Thr Asp Gly Lys Asp Lys Pro Asn His Gly Lys Ser Ile Thr Leu Lys Gly Ser Gly Thr Leu Thr Leu Asn Asn His Ile Asp Gln Gly Ala Gly Gly Leu Phe Phe Glu Gly Asp Tyr Glu Val Lys Gly Thr Ser Asp Ser Thr Thr Trp Lys Gly Ala Gly Val Ser Val Ala Asp Gly Lys Thr Val Thr Trp Lys Val His Asn Pro Lys Tyr Asp Arg Leu Ala Lys Ile Gly Lys Gly Thr Leu Val Val Glu Gly Lys Gly Lys Asn Glu Gly Leu Leu Lys Val Gly Asp Gly Thr Val Ile Leu Lys Gln Lys Ala Asp Ala Asn Asn Lys Val Gln Ala Phe Ser Gln Val Gly Ile Val Ser Gly Arg Ser Thr Leu Val 505 Leu Asn Asp Asp Lys Gln Val Asp Pro Asn Ser Ile Tyr Phe Gly Phe Arg Gly Gly Arg Leu Asp Leu Asn Gly Asn Ser Leu Thr Phe Asp His 535 Ile Arg Asn Ile Asp Asp Gly Ala Arg Val Val Asn His Asn Met Thr Asn Thr Ser Asn Ile Thr Ile Thr Gly Glu Ser Leu Ile Thr Asn Pro 565 570 Asn Thr Ile Thr Ser Tyr Asn Ile Glu Ala Gln Asp Asp Asp His Pro Leu Arg Ile Arg Ser Ile Pro Tyr Arg Gln Leu Tyr Phe Asn Gln Asp Asn Arg Ser Tyr Tyr Thr Leu Lys Lys Gly Ala Ser Thr Arg Ser Glu 615 Leu Pro Gln Asn Ser Gly Glu Ser Asn Glu Asn Trp Leu Tyr Met Gly 630 Arg Thr Ser Asp Ala Ala Lys Arg Asn Val Met Asn His Ile Asn Asn Glu Arg Met Asn Gly Phe Asn Gly Tyr Phe Gly Glu Glu Thr Lys Ala Thr Gln Asn Gly Lys Leu Asn Val Thr Phe Asn Gly Lys Ser Asp Gln Asn Arg Phe Leu Leu Thr Gly Gly Thr Asn Leu Asn Gly Asp Leu Asn Val Glu Lys Gly Thr Leu Phe Leu Ser Gly Arg Pro Thr Pro His

Ala Arg Asp Ile Ala Gly Ile Ser Ser Thr Lys Lys Asp Pro His Phe 730 Thr Glu Asn Asn Glu Val Val Val Glu Asp Asp Trp Ile Asn Arg Asn 745 Phe Lys Ala Thr Thr Met Asn Val Thr Gly Asn Ala Ser Leu Tyr Ser 760 Gly Arg Asn Val Ala Asn Ile Thr Ser Asn Ile Thr Ala Ser Asn Asn Ala Gln Val His Ile Gly Tyr Lys Thr Gly Asp Thr Val Cys Val Arg
785 790 795 800 Ser Asp Tyr Thr Gly Tyr Val Thr Cys His Asn Ser Asn Leu Ser Glu Lys Ala Leu Asn Ser Phe Asn Pro Thr Asn Leu Arg Gly Asn Val Asn 825 Leu Thr Glu Asn Ala Ser Phe Thr Leu Gly Lys Ala Asn Leu Phe Gly 840 Thr Ile Gln Ser Ile Gly Thr Ser Gln Val Asn Leu Lys Glu Asn Ser 855 His Trp His Leu Thr Gly Asn Ser Asn Val Asn Gln Leu Asn Leu Thr Asn Gly His Ile His Leu Asn Ala Gln Asn Asp Ala Asn Lys Val Thr Thr Tyr Asn Thr Leu Thr Val Asn Ser Leu Ser Gly Asn Gly Ser Phe 905 Tyr Tyr Trp Val Asp Phe Thr Asn Asn Lys Ser Asn Lys Val Val Val 920 Asn Lys Ser Ala Thr Gly Asn Phe Thr Leu Gln Val Ala Asp Lys Thr 935 Gly Glu Pro Asn His Asn Glu Leu Thr Leu Phe Asp Ala Ser Asn Ala Thr Arg Asn Asn Leu Glu Val Thr Leu Ala Asn Gly Ser Val Asp Arg Gly Ala Trp Lys Tyr Lys Leu Arg Asn Val Asn Gly Arg Tyr Asp Leu Tyr Asn Pro Glu Val Glu Lys Arg Asn Gln Thr Val Asp Thr Thr Asn 1000 Ile Thr Thr Pro Asn Asp Ile Gln Ala Asp Ala Pro Ser Ala Gln Ser 1015 1020 Asn Asn Glu Glu Ile Ala Arg Val Glu Thr Pro Val Pro Pro Pro Ala 1030 1035 Pro Ala Thr Glu Ser Ala Ile Ala Ser Glu Gln Pro Glu Thr Arg Pro 1045 1050 Ala Glu Thr Ala Gln Pro Ala Met Glu Glu Thr Asn Thr Ala Asn Ser 1065

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Thr Glu Thr Ala Pro Lys Ser Asp Thr Ala Thr Gln Thr Glu Asn Pro 1075 1080 1085

Asn Ser Glu Ser Val Pro Ser Glu Thr Thr Glu Lys Val Ala Glu Asn 1090 1095 1100

Pro Pro Gln Glu Asn Glu Thr Val Ala Lys Asn Glu Gln Glu Ala Thr 1105 1110 1115 1120

Glu Pro Thr Pro Gln Asn Gly Glu Val Ala Lys Glu Asp Gln Pro Thr 1125 1130 1135

Val Glu Ala Asn Thr Gln Thr Asn Glu Ala Thr Gln Ser Glu Gly Lys

Thr Glu Glu Thr Gln Thr Ala Glu Thr Lys Ser Glu Pro Thr Glu Ser

Val Thr Val Ser Glu Asn Gln Pro Glu Lys Thr Val Ser Gln Ser Thr 1170 1175 1180

Glu Asp Lys Val Val Val Glu Lys Glu Glu Lys Ala Lys Val Glu Thr 1185 1190 1195 1200

Glu Glu Thr Gln Lys Ala Pro Gln Val Thr Ser Lys Glu Pro Pro Lys 1205 1210 1215

Gln Ala Glu Pro Ala Pro Glu Glu Val Pro Thr Asp Thr Asn Ala Glu 1220 1225 1230

Glu Ala Gln Ala Leu Gln Gln Thr Gln Pro Thr Thr Val Ala Ala Ala 1235 1240 1245

Glu Thr Thr Ser Pro Asn Ser Lys Pro Ala Glu Glu Thr Gln Gln Pro 1250 1255 1260

Ser Glu Lys Thr Asn Ala Glu Pro Val Thr Pro Val Val Ser Glu Asn 1265 1270 1280

Thr Ala Thr Gln Pro Thr Glu Thr Glu Glu Thr Ala Lys Val Glu Lys 1285 1290 1295

Glu Lys Thr Gln Glu Val Pro Gln Val Ala Ser Gln Glu Ser Pro Lys 1300 1305 1310

Gln Glu Gln Pro Ala Ala Lys Pro Gln Ala Gln Thr Lys Pro Gln Ala 1315 1320 1325

Glu Pro Ala Arg Glu Asn Val Leu Thr Thr Lys Asn Val Gly Glu Pro 1330 1335 1340

Gln Pro Gln Ala Gln Pro Gln Thr Gln Ser Thr Ala Val Pro Thr Thr 1345 1350 1355 1360

Gly Glu Thr Ala Ala Asn Ser Lys Pro Ala Ala Lys Pro Gln Ala Gln 1365 1370 1375

Ala Lys Pro Gln Thr Glu Pro Ala Arg Glu Asn Val Ser Thr Val Asn 1380 1385 1390

Thr Lys Glu Pro Gln Ser Gln Thr Ser Ala Thr Val Ser Thr Glu Gln 1395 1400 1405

Pro Ala Lys Glu Thr Ser Ser Asn Val Glu Gln Pro Ala Pro Glu Asn 1410 1420

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Ser Ile Asn Thr Gly Ser Ala Thr Thr Met Thr Glu Thr Ala Glu Lys 1425 1430 1435 1440

- Ser Asp Lys Pro Gln Met Glu Thr Val Thr Glu Asn Asp Arg Gln Pro 1445 1450 1455
- Glu Ala Asn Thr Val Ala Asp Asn Ser Val Ala Asn Asn Ser Glu Ser 1460 1465 1470
- Ser Glu Ser Lys Ser Arg Arg Arg Arg Ser Val Ser Gln Pro Lys Glu 1475 1480 1485
- Thr Ser Ala Glu Glu Thr Thr Val Ala Ser Thr Gln Glu Thr Thr Val 1490 1495 1500
- Asp Asn Ser Val Ser Thr Pro Lys Pro Arg Ser Arg Arg Thr Arg Arg 1505 1510 1515 1520
- Ser Val Gln Thr Asn Ser Tyr Glu Pro Val Glu Leu Pro Thr Glu Asn 1525 1530 1535
- Ala Glu Asn Ala Glu Asn Val Gln Ser Gly Asn Asn Val Ala Asn Ser 1540 1545 1550
- Gln Pro Ala Leu Arg Asn Leu Thr Ser Lys Asn Thr Asn Ala Val Ile 1555 1560 1565
- Ser Asn Ala Met Ala Lys Ala Gln Phe Val Ala Leu Asn Val Gly Lys 1570 1580
- Ala Val Ser Gln His Ile Ser Gln Leu Glu Met Asn Asn Glu Gly Gln 1585 1590 1595 1600
- Tyr Asn Val Trp Ile Ser Asn Thr Ser Met Asn Lys Asn Tyr Ser Ser 1605 1610 1615
- Glu Gln Tyr Arg Arg Phe Ser Ser Lys Ser Thr Gln Thr Gln Leu Gly
  1620 1625 1630
- Trp Asp Gln Thr Ile Ser Asn Asn Val Gln Leu Gly Gly Val Phe Thr 1635 1640 1645
- Tyr Val Arg Asn Ser Asn Asn Phe Asp Lys Ala Ser Ser Lys Asn Thr 1650 1655 1660
- Leu Ala Gln Val Asn Phe Tyr Ser Lys Tyr Tyr Ala Asp Asn His Trp 1665 1670 1680
- Tyr Leu Gly Ile Asp Leu Gly Tyr Gly Lys Phe Gln Ser Asn Leu Gln 1685 1690 1695
- Thr Asn Asn Ala Lys Phe Ala Arg His Thr Ala Gln Ile Gly Leu 1700 1705 1710
- Thr Ala Gly Lys Ala Phe Asn Leu Gly Asn Phe Ala Val Lys Pro Thr 1715 1720 1725
- Val Gly Val Arg Tyr Ser Tyr Leu Ser Asn Ala Asp Phe Ala Leu Ala 1730 1735 1740
- Gln Asp Arg Ile Lys Val Asn Pro Ile Ser Val Lys Thr Ala Phe Ala 1745 1750 1755 1760
- Gln Val Asp Leu Ser Tyr Thr Tyr His Leu Gly Glu Phe Ser Ile Thr 1765 1770 1775

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Pro Ile Leu Ser Ala Arg Tyr Asp Ala Asn Gln Gly Asn Gly Lys Ile 1785

Asn Val Ser Val Tyr Asp Phe Ala Tyr Asn Val Glu Asn Gln Gln 1800 1805

Tyr Asn Ala Gly Leu Lys Leu Lys Tyr His Asn Val Lys Leu Ser Leu 1815 1820

Ile Gly Gly Leu Thr Lys Ala Lys Gln Ala Glu Lys Gln Lys Thr Ala 1835

Glu Val Lys Leu Ser Phe Ser Phe 1845

- (2) INFORMATION FOR SEQ ID NO:7:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 8 amino acids
    - (B) TYPE: amino acid (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:7:
  - Gly Asp Ser Gly Ser Pro Met Phe 5
- (2) INFORMATION FOR SEQ ID NO:8:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 8 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:8:
  - Gly Asp Ser Gly Ser Pro Leu Phe
- (2) INFORMATION FOR SEQ ID NO:9:
  - (i) SEQUENCE CHARACTERISTICS:
    - (A) LENGTH: 7 amino acids
    - (B) TYPE: amino acid
    - (D) TOPOLOGY: linear
  - (xi) SEQUENCE DESCRIPTION: SEQ ID NO:9:

His Thr Tyr Phe Gly Ile Asp 1

25

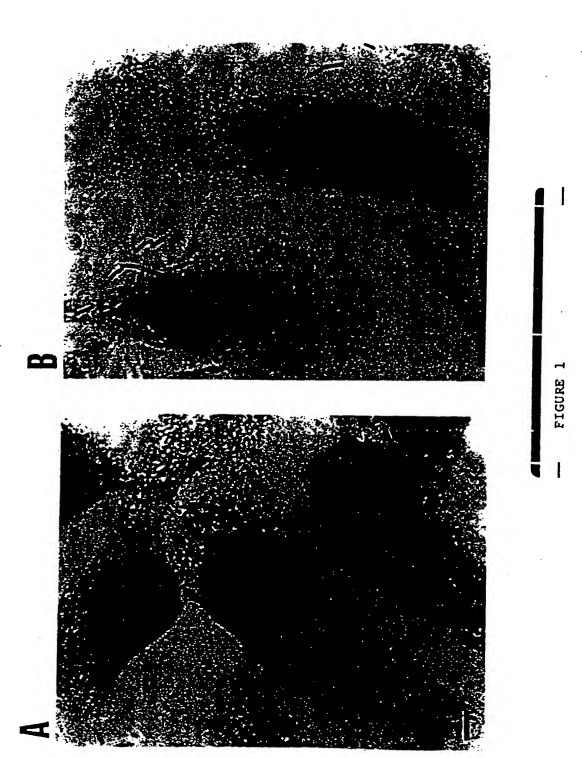
#### CLAIMS

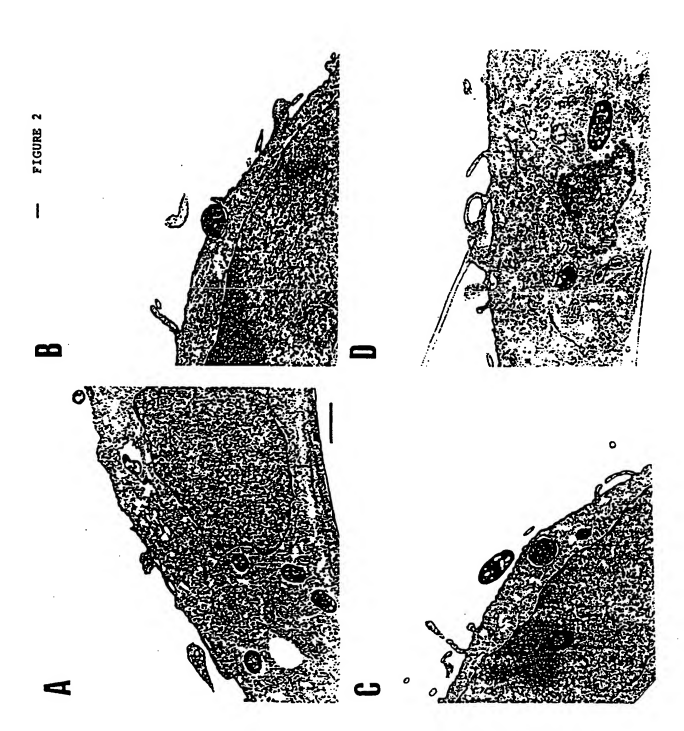
- 1. A recombinant Haemophilus adhesion and penetration protein.
- A recombinant Haemophilus adhesion and penetration
   protein according to claim 1 which has a sequence homologous to that shown in Figure 6.
  - 3. A recombinant *Haemophilus* adhesion and penetration protein according to claim 1 which has the sequence shown in Figure 6.
- 4. A recombinant nucleic acid encoding an Haemophilus adhesion and penetration protein.
  - 5. The nucleic acid of claim 3 comprising DNA having a sequence homologous to that shown in Figure 6.
- 6. An expression vector comprising transcriptional and translational regulatory nucleic acid operably linked to nucleic acid encoding an *Haemophilus* adhesion and penetration protein.
- A host cell transformed with an expression vector comprising a nucleic acid encoding an Haemophilus
   adhesion and penetration protein.
  - 8. A method of producing an Haemophilus adhesion and penetration protein comprising:
  - a) culturing a host cell transformed with an expressing vector comprising a nucleic acid encoding an Haemophilus adhesion and penetration protein; and

- b) expressing said nucleic acid to produce an Haemophilus adhesion and penetration protein.
- 9. A vaccine comprising a pharmaceutically acceptable carrier and an *Haemophilus* adhesion and penetration protein for prophylactic or therapeutic use in generating an immune response.
- 10. A vaccine according to claim 8 wherein said Haemophilus adhesion and penetration protein has a sequence homologous to that shown in Figure 6.
- 10 11. A monoclonal antibody capable of binding to an Haemophilus adhesion and penetration protein.

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12. A method of treating or preventing *Haemophilus* influenzae infection comprising administering the vaccine of claim 9 or 10.





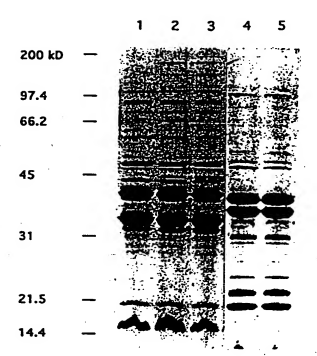
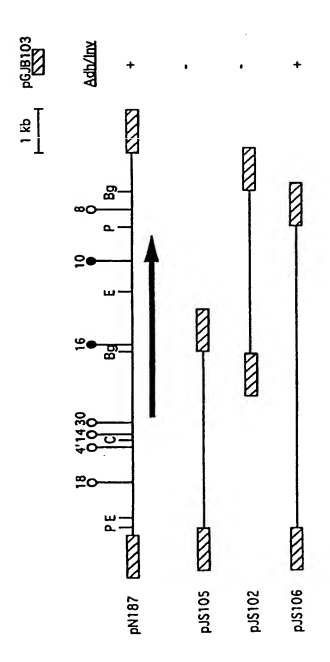


FIGURE 3





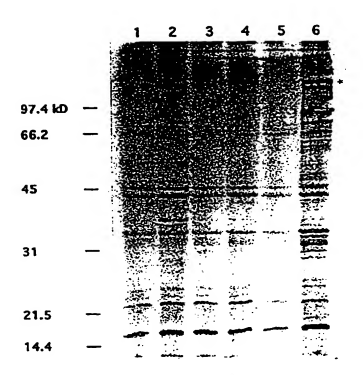


FIGURE 5

- MKKT -10 170 230 A E N K G K F T V G A Q N I K V Y N K Q G Q L V G T S M T K GCCCCGATGATTGATTTTTCTGTAGTGTCACGTAACGGCGTGGCAGCCTTGGTTGAAAATCAATATATTGTGAGCGTGGCACATAACGTA 310 A P M I D F S V V S R N G V A A L V E N Q Y I V S V A H N V GGATATACAGATGTTGATTTTGGTGCAGAGGGAAACAACCCCGATCAACATCGTTTTACTTATAAGATTGTAAAACGAAATAACTACAAA
  G Y T D V D F G A E G N N P D Q H R F T Y K I V K R N N Y K 530 510 AAAGATAATTTACATCCTTATGAGGACGATTACCATAATCCACGATTACATAAATTCGTTACAGAAGCGGCTCCAATTGATATGACTTCG K D N L H P Y E D D Y H N P R L H K F V T E A A P I D M T S AATATGAATGGCAGTACTTATTCAGATAGAACAAAATATCCAGAACGTGTTCGTATCGGCTCTGGACGGCAGTTTTGGCGAAATGATCAA N N G S T Y S D R T K Y P E R V R I G S G R Q F W R N D Q 690 670 GACAAAGGCGACCAAGTTGCCGGTGCATATCATTATCTGACAGCTGGCAATACACACAATCAGCGTGGAGCAGGTAATGGATATTCGTAT D K G D Q V A G A Y H Y L T A G N T H N Q R G A G N G Y S Y 790 770 750 GATGCT GAAAAACAAAAAT GGTTAATTAAT GGGATATTAC GGGAAGGCAACCCCTTTT GAAGGCAAAGAAAAT GGGTTTCAATT GGTT CGC DAEKQKWLINGILREGNPFEGKENGFQLVR AAATCTTÄTTTTGATGAAATTTTCGAAAGAGATTTACATCACTTTACACCCGAGCTGGTAATGGAGTGTACACAATTAGTGGAAAT K S Y F D E I F E R D L H T S L Y T R A G 'N G V Y T I S G H 1050 1030 GATAAT GGTCAGGGGTCTATAACTCAGAAATCAGGAATACCATCAGAAATTAAAATTACGTTAGCAAATATGAGTTTACCTTTGAAAGAG D N G Q G S I T Q K S G I P S E I K I T L A N M S L P L K E

  - 1270 1290 1310 1330 1350

    AACCAAACTTGGCAAGGAGCTGGCATACATGTAAGTGAAAATAGCACCGTTACTTGGAAAGTAAATGGCGTGGAACATGATCGACTTTCT

    N Q T W Q G A G I H V S E N S T V T W K V N G V E H D R L S
  - 1370

    AAAATTGGTAAAGGAACATTGCACGTTCAAGCCAAAGGGGAAAATAAAGGTTCGATCAGCGTAGGCGATGGTAAAGTCATTTTGGAGCAG
    K I G K G T L H V Q A K G E H K G S I S V G D G K V I L E Q

FIGURE 6A

1450 1470 1490 1510 CAGGCAGACGATCAAGGCAACAAACAAGCCTTTAGTGAAATTGGCTTGGTTAGCGGCAGAGGGACTGTTCAATTAAACGATGATAAACAA Q A D D Q G N K Q A F S E I G L V S G R G T V Q L N D D K Q 1570 1590 FDTDKFYFGFRGGRLDLNGHSLTFKRIQNT GACGAGGGGCAATGATTGTGAACCATAATACAACTCAAGCCGCTAATGTCACTATTACTGGGAACGAAAGCATTGTTCTACCTAATGGA D E G A M I V N H N T T Q A A N V T I T G N E S I V L P N G 1690 N N I N K L D Y R K E I A Y N G W F G E T D K N K H N G R L 1830 1850 1930 K L F F S G R P T P H A Y N H L N K R W S E M E G I P Q G E 2030 ATTGTGTGGGATCACGATTGGATCAACCGTACATTTAAAGCTGAAAACTTCCAAATTAAAGGCGGAAGTGCGGTGGTTTCTCGCAATGTT IV W. D H D W I N R T F K A E N F Q I K G G S A V V S R N V TCTTCAATTGAGGGAAATTGGACAGTCAGCAATAATGCAAATGCCACATTTGGTGTTGTGCCAAATCAACAAAATACCATTTGCACGCGT 2130 SSIEGNWTVSHNAHATFGVVPHQQNTICTR 2190 TCAGATTGGACAGGATTAACGACTTGTCAAAAAGTGGATTTAACCGATACAAAAGTTATTAATTCTATACCAAAAACACAAATCAATGGC 2210 S D W T G L T T C Q K V D L T D T K V I N S I P K T Q I N G 2290 TCTATTAATTTAACTGATAATGCAACGGCGAATGTTAAAGGTTTAGCAAAACTTAATGGCAATGTCACTTTAACAAATCACAGCCAATTT
5 I N L T D N A T A N V K G L A K L N G N V T L T N H S Q F 2310 2370 2390 ACATTAAGCAACAATGCCACCCAAATAGGCAATATTCGACTTTCCGACAATTCAACTGCAACGGTGGATAATGCAAACTTGAACGGTAAT T L S N N A T Q I G N I R L S D N S T A T V D N A N L N G N 2470 **2490** GTGCATTTAACGGATTCAGCTCAATTTTCTTTAAAAAACAGCCATTTTTCGCACCAAATTCAGGGAGACAAAGGCACAACAGTGACGTTG V H L T D S A Q F S L K N S H F S H Q I Q G D K G T T V T L 2570 GAAAAT GCGACTT GGACAAT GCCTAGCGATACTACATT GCAGAATTTAACGCTAAATAACAGTACGATCACGTTAAATTCAGCTTATTCA 2590 ENATWIM PSDTTLQNLTLNNSTITLNSAYS 2650 GCTAGCTCAAACAATACGCCACGTCGCCGTTCATTAGAGACGGAAACAACGCCAACATCGGCAGAACATCGTTTCAACACATTGACAGTA 2670 ASSNNTPRRRSLETETTPTSAEHRFNTLT 2730 2750 AATGGTAAATTGAGTGGGCAAGGCACATTCCAATTTACTTCATCTTTATTTGGCTATAAAAGCGATAAATTAAAATTATCCAATGACGCT N G K L S G Q G T F Q F T S S L F G Y K S D K L K L S N D A 2770 GAGGGCGATTACATATTATCTGTTCGCAACACAGGCAAAGAACCCGAAACCCTTGAGCAATTAACTTTGGTTGAAAGCAAAGATAATCAA E G D Y I L S V R N T G K E P E T L E Q L T L V E S K D N Q

FIGURE 6B

- 2890 2910 2930 2950 2970
  CCGTTATCAGATAAGCTCAAATTTACTTTAGAAAATGACCACGTTGATGCAGGTGCATTACGTTATAAATTAGTGAAGAATGATGGCGAA
  P L S D K L K F T L E N D H V D A G A L R Y K L V K N D G E

- 3250 3270 3290 3310 3330

  AAAAGAGCCAGTGTTTCTGATCCCCTGCTTGATCAAAGCCTGTTCGCATTAGAAGCCGCACTTGAGGTTATTGATGCCCCACAGCAATCG
  K R A V F S D P L L D Q S L F A L E A A L E V I D A P Q Q S
- 3430
  3450
  3470
  3490
  3510
  TTATCTGCAACAGTAAATAGTATGCTTTCTGTTCAAGATGAATTAGATCGTCTTTTTGTAGATCAAGCACAATCTGCCGTGTGGACAAAT
  L S A T V N S N L S V Q D E L D R L F V D Q A Q S A V W T N
- 3530 3550 3570 3590
  ATCGCACAGGATAAAAGACGCTATGATTCTGATGCGTTCCGTGCTTATCAGCAGCAGAAAACGAACTTACGTCAAATTGGGGTGCAAAAA
  I A Q D K R R Y D S D A F R Å Y Q Q Q K T N L R Q I G V Q K
- 3610
  3630
  3650
  3670
  3690
  GCCTTAGCTAATGGACGAATTGGGGCAGTTTCTCGCATAGCCGTTCAGATAATACCTTTGATGAACAGGTTAAAAATCACGCGACATTA
  A L A N G R I G A V F S H S R S D N T F D E Q V K N H A T L
- 3730 3750 3770

  ACGATGATGTCGGGTTTTGCCCAATATCAATGGGGCGATTTACAATTTGGTGTAAACGTGGGAACGGGAATCAGTGCGAGTAAAATGGCT
  T M M S G F A Q Y Q W G D L Q F G V N V G T G I S A S K M A
- 3790 3810 3830 3850 3870

  GAAGAACAAAGCCGAAAAATTCATCGAAAAGCGATAAATTATGGCGTGAATGCAAGTTATCAGTTCCGTTTAGGGCAATTGGGCATTCAG
  E E Q S R K I H R K A I N Y G V N A S Y Q F R L G Q L G I Q
- 3890 3910 3930 3950
  CCTTATTTTGGAGTTAATCGCTATTTTATTGAACGTGAAAATTATCAATCTGAGGAAGTGAGAGTGAAAACGCCTAGCCTTGCATTTAAT
  PYFGVNRYFIERENYQSEEVRVKTPSLAFN
- 3970 3990 4010 4030 4050 CGCTATAATGCTGGGGTTAATGCTGGATTATATATGTTGAT R Y N A G I R V D Y T F T P T D N I S V K P Y F F V N Y V D
- 4070 4090 4110 4130
  GTTTCAAACGCTAACGTACAACCACGGTAAATCTCACGGTGTTGCAACAACCATTTGGACGTTATTGGCAAAAAGAAGTGGGATTAAAG
  V S N A N V Q T T V N L T V L Q Q P F G R Y W Q K E V G L K
- 4150 4170 4190 4210 4230

  GCAGAAATTTTACATTTCCAAATTTCCGCTTTTATCTCAAAATCTCAAGGTTCACAACTCGGCAAACAGCAAAATGTGGGCGTGAAATTG
  A E I L H F Q I S A F I S K S Q G S Q L G K Q Q N V G V K L
- 4250 4270 4290 4310
  GGCTATCGTTGGTAAAAATCAACATAATTTTATCGTTTATTGATAAACAAGGTGGGTCAGATCAGATCCCACCTTTTTTATTCCAATAAT

- FIGURE 6C

Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1 MKKTVFRINE LTACISLGIV MINKKEKINE IALTVAYALT MINKKEKINE IALTVAYALT MINKKEKINE IALTVAYALT MINKKEKINE IALTVAYALT MINKKEKINE IALTVAYALT	PYTEAALVRD PYTEAALVRD PYTEAALVRD PYTEAALVRD	DVDYQIFFOF DVDYQIFFOF DVDYQIFFOF	AENKGKESVG AENKGKESVG AENKGKESVG AENKGKESVG
Hap HK368IGA HK393IG HK715IGA HK61IGA Consensus	51 AQNIKVYNKQ GQLVGTSMIK ATNVLVKDKN NKDLGTALPN ATNVEVROKN NRPLGNVLPN ATNVEVROKN NHSLGNVLPN ATNVEVROKK NQSLGSALPN A-NVKG	GIPMIDESVV GIPMIDESVV GIPMIDESVV	DVDKRIATLI DVDKRIATLI DVDKRIATLI DVDKRIATLV	NEGANACAKH NEGANACAKH NEGANACAKH
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	101NVGY TDVDFGAEGN VSNGVSELHF GNINGMNNG VSNGVSELHF GNINGMNNG VSNGVSELHF GNINGMNNG VSNGVSELHF GNINGMNNG	NAKAHROVSS NAKAHROVSS NDKSHROVSS NAKSHROVSS	EENRYFSVEK EENRYYTVEK EENRYFSVEK EENRYYTVEK	NEYPTKLNGK NEYPTKLNGK
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	151KKONLH PYEDDYHNPR TVTTEDQ.TQ KRREDYYMPR AVITEDQ.AQ KRREDYYMPR AVITEDQ.TQ KRREDYYMPR FTTKEEQDAQ KRREDYYMPR	LDKFVTEVAP LDKFVTEVAP LDKFVTEVAP	IEASTASSDA IEASTASSDA IEASTANNIK	GTYNDQNKYP GTYNDQNKYP GEYNNSDKYP
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	201 ERVRIGSGRO F AFVRLGSGSQ FIYKKEDNYS YFVRLGSGSQ FIYKKEDNYS AFVRLGSGSQ FIYKKEDNYS AFVRLGSGSQ FIYKKGSRYQVR-GSG-Q F	LILN LULG LILN LILTEKDKQG	NHEVGG KEGQKSDAGG NHEVGG NLLRNWDVGG	NNLKLVGDAY NNLKLVGDAY DNLELVGNAY

Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	251 300 HYLTAGNIHN QRGAGNGYSY LGGD VRKAGEYGPL PIAGSKGDSG TYGIAGIPYK VNHENNGLIG FGNSKEEHSD PKGILSQDPL TNYAVLGDSG TYGIAGIPYE VNHENNGLIG FGNSKEEHSD PKGILSQDPL TNYAVLGDSG TYGIAGIPYK VNHENNGLIG FGNSKEEHSD PKGILSQDPL TNYAVLGDSG TYGIAGIPYK VNHENNGLIG FGNSKEEHSD PKGILSQDPL TNYAVLGDSG TYGIAGIPYK VNHENNGLIG FGNSKEEHSD PKGILSQDPL TNYAVLGDSG -YAG
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	350 SPMFTYDAEK OKWILINGILR EGNPFECKEN GFOLVRKSYF D.EIFERDLH SPIFVYDREK GKWIFIGSYD FWAGYNKKSWQ EWNIYKSOFT SPIFVYDREK GKWIFIGSYD YWAGYNKKSWQ EWNIYKPEFA SPIFVYDREK GKWIFIGSYD FWAGYNKKSWQ EWNIYKPEFA SPIFVYDREK GKWIFIGSYD FWAGYNKKSWQ EWNIYKHEFA SP-F-YD-EK -KWIG
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	351  TSLYTRAGNG VYTISGNDNG QGSITQKSGI PSEIKITLAN MSLPLKEKOK KDVLNKDSAG SLIGSKTDYS WSSNGKTSTI TGGEKS INVDLAD EKIYEQYSAG SLIGSKTDYS WSSNGKTSTI TGGEKS INVDLAD KTVLDKDTAG SLTGSNTQYN WNPTGKTSVI SNGSES INVDLFD EKIYQQYSAG SLTGSNTQYT WQATGSTSTI TGGGEP LSVDLTD
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	450 VHNPRYDGPN IYSPRINNGE TLYFMDOKQG SLIFASDINQ GAGGLYFEGN
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	451 500 FTVSPNSNQ. TWQGAGIHVS ENSTVTWKVN GVEHDRLSKI GKGTLHVQAK YEVKGTSDNT TWKGAGVSVA EGKTVTWKVH NPQYDRLAKI GKGTLIVEGT YEVKGTSDNT TWKGAGVSVA EGKTVTWKVH NPQYDRLAKI GKGTLIVEGT YEVKGTSDST TWKGAGVSVA DGKTVTWKVH NPKSDRLAKI GKGTLIVEGK YEVKGTSDST TWKGAGVSVA DGKTVTWKVH NPKYDRLAKI GKGTLIVEGK YEVKGTSDST TWKGAGVSVA DGKTVTWKVH NPKYDRLAKI GKGTLIVEGK

Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	501 GENKGSISVG DGKVI GDNKGSIKVG DGTVI GENKGSIKVG DGTVI GENKGSIKVG DGTVI GKNEGILKVG DGTVI G-N-GVG DG-VI	IKOOT NGSGO.HAFA IKOOT NGSGO.HAFA IKOOA DANNKVKAFS IKOKA DANNKVOAFS	SVGIVSGRST SVGIVSGRST QVGIVSGRST QVGIVSGRST	TATINDDKÖAD AATINDDKÖAD TATINDDKÖAD TATINDDKÖAD
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	551 TDKFYFGERG GRLDLI PNSIYFGERG GRLDLI PNSIYFGERG GRLDLI PNSIYFGERG GRLDLI PNSIYFGERG GRLDLIYFGERG GRLD-	NGNSL TFDHIRNIDD NGNSL TFDHIRNIDD NGNNL TFEHIRNIDD NGNSL TFDHIRNIDD	GARLVNHIMT GARLVNHISTS GARLVNHINTS GARVVNHIMT	NASNITITGE KHSTVTITGD KTSTVTITGE NTSNITITGE
-	601			 650
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	ESIVLPNG SLITDPNTIT PYNID NLITDPNNVS IYYVK SLITDPNTIT PYNID SLITNPNTIT SYNIEIPN	PLEDD NPYATRQIKY APDED NPYAFRRIKD AQDDD HPLRIRSIPY	GCOLYINLEN GYOLYFNEEN GCOLYINLEN	RTYYALKKDA YTYYALRKGA
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	STRSELPKNS GESNE SIRSEFPONR GESNN STRSELPKNS GESNE STRSELPONS GESNE	ISWLYM GTEKADAOKN NWLYM GKTSDEAKRN NWLYM GRTSDEAKRN	VMNHINNERM AMNHINNERM VMNHINNERM	NGFNGYFGEE NGFNGYFGEE NGFNGYFGEE NGFNGYFGEE
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	701 D.KNKHNGRL NLIYK EGKNNGNL NVIFK EGKNNGNL NVIFK EGKNNGNL NVIFK ETKATONGKL NVIFNKNG-L N	CCKSEQ NRFILITGGIN CCKSEQ NRFILITGGIN CCKSEQ NRFILITGGIN	INCOLIVEKG INCOLIVEKG INCOLIVEKG	TLFLSGRPTP TLFLSGRPTP TLFLSGRPTP TLFLSGRPTP

## 12/19:

Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	751 HAYNHINKRW SEMEGIPQ GETVADHDWI NRTFKAENEQ IKGGSAVVS. HARDIAGISS TKKDPHFAEN NEVVVEDDWI NRNFKATIMN VIGNASLYSG HARDIAGISS TKKDSHFSEN NEVVVEDDWI NRNFKATNIN VINNATLYSG HARDIAGISS TKKDOHFAEN NEVVVEDDWI NRNFKATIMN VIGNASLYSG HARDIAGISS TKKDPHFTEN NEVVVEDDWI NRNFKATIMN VIGNASLYSG HAR
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	RNVSSIEGW TVSNIANATE GVVPNQNTI CTRSDWTGLT TCQKVDLTDT RNVANITSNI TASNKAQVHI GYKTGDTV CVRSDYTGYV TCTTDKLSD. RNVANITSNI TASNKAKHI GYKAGDTV CVRSDYTGYV TCTTDKLSD. RNVANITSNI TASNKAVHI GYKAGDTV CVRSDYTGYV TCTTDKLSD. RNVANITSNI TASNKAQVHI GYKTGDTV CVRSDYTGYV TCHNSNLSE. RNVANITSNI TASNKAQVHI GYKTGDTV CVRSDYTGYV TCHNSNLSE. RNV-I-N- T-S-A GT- C-RSD-TG TCL
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	851 KVINSIPKTO INGSINLTDN ATANVKGLAK INGNVTLINH SOFTLSNNAT KALNSFNPTN LRGNVNLTES A KALNSFNATN VSGNVNLSGN A KALNSFNPTN LRGNVNLTEN A KALNSFNPTN LRGNVNLTEN A  K-NSTGNL A
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	901 QIGNIRLSDN STATVDNANL NGNVHLIDSA QFSLKNSHFS HQIQQDKGTT .NFVLGKANL FGTIQSRGNS QVRLT
Hæp HK368IGA HK393IGA HK715IGA HK61IGA Consensus	951 VTLENATWIM PSDTTLQNLT LINISTITLIS AYSASSINIP RRRSLETETTENSHWHL TQNSDVHQLD LANGHIHLINS ADNSNIVTKENSHWHL TQNSDVHQLD LANGHIHLINS ADNSNIVTKENSHWHL TQDSNIVQLN LDKGHIHLINA QNDANKVITENSHWHL TQNSNIVQLN LTNGHIHLINA QNDANKVITENSHWHL TQNSNIVQLN LTNGHIHLINA QNDANKVIT.

Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1050 PTSAEHRENT LTVNCKLSGQ GTFQFTSSLF GYKSDKLKLS NDAEGDYILSYNT LTVNS.LSGN GSFYYLTDLS NKQGDKVVVT KSATGNFTLQYNT LTVNS.LSGN GSFYYLTDLS NKQGDKVVVT KSATGNFTLQYNT LTVNS.LSGN GSFYYLTDLS NKQGDKVVVT KSATGNFTLQYNT LTVNS.LSGN GSFYYWDFT NNKSNKVVVN KSATGNFTLQNT LTVN-LSG- G-F
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1100 VRNICKEPET LEQLTLVESK DNOPLSDKLK FTLENDHVDA GALRYKLVKN VADKTGEPNH NELTLFDAS KAOR. DHLN VSLVGNIVDL GAWKYKLRNV VADKTGEPTK NELTLFDAS NATR. NNLN VSLVGNIVDL GAWKYKLRNV VADKTGEPH NELTLFDAS NATR. NNLN VSLVGNIVDL GAWKYKLRNV VADKTGEPH NELTLFDAS NATR. NNLE VTLANGSVDR GAWKYKLRNV VEPLTLVD- GAYKL
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1101 DGEFRLHNPI KEQELHNDLV NGRYDLYNPEVEKRNQIV DTINITIPNN IQADVPSVPS NNEEIARVDE NGRYDLYNPEVEKRNQIV DTINITIPNN IQADVPSVPS NNEEIARVDE NGRYDLYNPEVEKRNQIV DTINITIPNN IQADVPSVPS NNEEIARV.E NGRYDLYNPEVEKRNQIV DTINITIPND IQADAPSAQS NNEEIARV.E -GL-NPE-ENV
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1151 1200 APVPPPAPAT
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1250

Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1251AKTQT GE  KSNVKANTQT NEVAQSGSET KETQTTETKETATVE KSNVKANTQT NEVAQSGSET KETQTTETKETATVE KPSVKANTQT NEVAQSGSET KETQTTETKETAKVE CPTVEANTQT NEATQSEGKT EETQTAETKS EPTESVTVSE NOPEKTVSQSA-TQT -E
Hap HK368IGA HK393IGA	1301 1350  KEEK
HK715IGA HK61IGA Consensus	TEDKVVVEKE EKAKVETEET QKAPQVTSKE PPAQAEPAPE EVI IDITUELO
Hap HK368ICA HK393IGA HK715IGA HK61IGA Consensus	TOVOAOPOTO STIVAAAEAT SPNSKPAEET .OPSEKTNAE PVIPVVSKNO A. QALQOTO PTIVAAAETT SPNSKPAEET QOPSEKTNAE PVIPVVS
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1450
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1451 1500

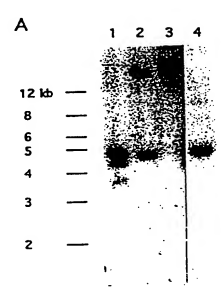
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1500 D QSLINALFAKQAEL TAETQKSKAK TKK  QPQAEPAREN DPTVNIKEPQSQTNT TADTEQPAKE TSSNVE  QPQAEPAREN DPTVNIKEPQSQTNT TADTEQPAKE TSSNVE  QPQAVLESEN VPTVNNAEEV QAQLQTQTSA TVSTKQPAPE NSINIG  KPQTEPAREN VSTVNIKEPQSQTSA TVSTEQPAKE TSSNVEQPAP
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1551 1600  V RSKRAVFSDP LLDQSL  OPVT ESTTVNTQNS VVEN  OPVT ESTTVNTQNS VVEN  SAT ALTETAEKSD KPQTETAAST EDASQHKANT VADNSVANNS ENSINTGSAT TMTETAEKSD KPQMETVT ENDRQPEANT VADNSVANNS
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1650
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1700QKDLI SRYSNSALSE SVRSVPHNVE PATTSSNDRSTVALCDLT STNTNAVLSD SVRSVPHNVE PATTSSNDRSTVALCDLT STNTNAVLSD SVRSE PTVINGSDRSTVALRDLT STNTNAVISD SVQTNSYEPV ELPTENAENA ENVOSQNVA NSOPALRNLT SKNTNAVLSN
Hap HK3681CA HK3931CA HK7151CA HK611CA Consensus	1750 LSATV NSMLSVQDEL DRL.FVDQAQ SAVWINIAQD KRRYDSDAFR ARAKAQFVAL NVGKAVSQHI SQLEMNEGQ YNVWVSNTSM NKNYSSSQYR ARAKAQFVAL NVGKAVSQHI SQLEMNEGQ YNVWSNTSM NENYSSSQYR AMAKAQFVAL NVGKAVSQHI SQLEMNEGQ YNVWSNTSM NENYSSSQYR AMAKAQFVAL NVGKAVSQHI SQLEMNEGQ YNVWISNTSM NKNYSSEQYRA NVLQVWY-SR

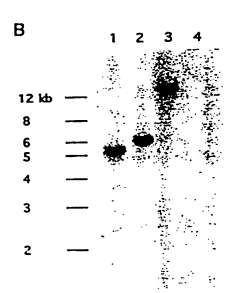
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1800 1751 AYQQQKTNLR QIGVQKALAN GRIGAVESHS RSDNTFDEQV KNHATLTMMS. RESSKSTQTQ LGWDQTISNN VQLGGVETYV RNSNNEDKAT SKN.TLAQVN RESSKSTQTQ LGWDQTISNN VQLGGVETYV RNSNNEDKAS SKN.TLAQVN
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	GFAQYQWGDL OFGVNVGT GISASKMAEE QSRKIHRKAI NYGVNASYQF FYSKY.YADN HWYLGIDLGY GKFQSKLQTN HNAKFARHTA OFGLTAGKAF FYSKY.YADN HWYLGIDLGY GKFQSNLXTN HNAKFARHTA OFGLTAGKAF FYSKY.YADN HWYLGIDLGY GKFQSNLXTN HNAKFARHTA OFGLTAGKAF FYSKY.YADN HWYLGIDLGY GKFQSNLQTN NNAKFARHTA QIGLTAGKAF FYSKY.YADN HWYLGIDLGY GKFQSNLQTN NNAKFARHTA QIGLTAGKAF YDGG- GSKRGAF
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1851 RLGQLGIQPY FGVNRYFIER ENYOSEEVRV KTPSLAFNRY NAGIRVDYTF NLGNFGITPI VGVRYSYLSN ADFALDQARI KVNPISVKTA FAQVDLSYTY NLGNFGITPI VGVRYSYLSN ANFALAKDRI KVNPISVKTA FAQVDLSYTY NLGNFAVKPT VGVRYSYLSN ADFALAQDRI KVNPISVKTA FAQVDLSYTY NLGNFAVKPT VGVRYSYLSN ADFALAQDRI KVNPISVKTA FAQVDLSYTY -LGPGV
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1901 TPTDNISVKP YFFVNYVDVS NANVQTTVNL TVLQQPFGRY WQKEVGLKAE .HLGEFSVTP ILSARY.DAN QGSGKINVNG YDFAYNVENQ QQYNAGLKLK .HLGEFSVTP ILSARY.DIN QGSGKINVNQ YDFAYNVENQ QQYNAGLKLK .HLGEFSVTP ILSARY.DIN QGSGKINVNQ YDFAYNVENQ QQYNAGLKLK .HLGEFSITP ILSARY.DAN QGNGKINVSV YDFAYNVENQ QQYNAGLKLKSPY-DVQGLK
Hap HK368IGA HK393IGA HK715IGA HK61IGA Consensus	1951 11HFQISAFI SKSQGSQLGK QQNVGVKLGY RW YHNVKLSLIG GLTKAKQAEK QKTAELKLSF SF YHNVKLSLIG GLTKAKQAEK QKTAELKLSF SF YHNVKLSLIG GLTKAKQAEK QKTAEVKLSF SF YHNVKLSLIG GLTKAKQAEK QKTAEVKLSF SF

1 2 3

105.1 kD —
69.8 —
43.3 —
28.3 —

- FIGURE 8





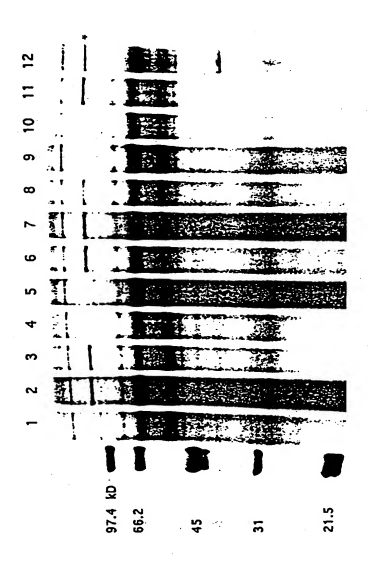


FIGURE 10

#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US95/10661

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A. CLASSIFICATION OF SUBJECT MATTER  IPC(6) : Please See Extra Sheet.			
US CL	:Please See Extra Sheet. to International Patent Classification (IPC) or to both	national classification and IPC	
	LDS SEARCHED		
	locumentation searched (classification system follower	d by classification symbols)	
			0/250 387 1 388 1
	424/130.1, 139.1, 150.1, 164.1, 184.1, 185.1, 242.		
Documentat	tion searched other than minimum documentation to the	e extent that such documents are included	in the fields searched
Electronic d	lata base consulted during the international search (na	ime of data base and, where practicable	, search terms used) .
·			
C. DOC	UMENTS CONSIDERED TO BE RELEVANT		
Category*	Citation of document, with indication, where ar	propriate, of the relevant passages	Relevant to claim No.
X	Infection and Immunity, Volume		
	1992, Barenkamp et al, "Clonin		12
Y		Encoding Nontypeable	12
	Haemophilus influenzae High-M		
	Exposed Proteins Related to Filan		
	Bordetella pertussis", pages 1302	2-1313, see pages 1302,	
	1303,1310, 1312, see Abstract.	A.	
X	Infection and Immunity, Volume 1990, Thomas et al, "Expression High-Molecular-Weight Protective Nontypeable and Type b Haemo 1909-1913, see pages 1909, 191	n in Escherichia coli of a Surface Antigen Found in philus influenzae", pages	1-8, 11
X Furth	ner documents are listed in the continuation of Box C	. See patent family annex.	
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19 OCTOBER 1995 28 NOV 1995			
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International application No.
PCT/US95/10661

Category*	Citation of document, with indication, where appropriate, of the relevant passages	Palamaria	
K	Proceedings of the National Academy of Sciences, Volume 90, issued April 1993 Geme III et al, "High-molecular-weight proteins of nontypable Haemophilus influenzae mediate attachment to human epithelial cells", pages 2875-2879, see pages 2875, 2876.	Relevant to claim No	
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#### INTERNATIONAL SEARCH REPORT

International application No. PCT/US95/10661

A. CLASSIFICATION	OF SUBJECT	MATTER
IDC (6).		

A61K 39/00, 39/02, 39/40, 39/102, 39/395; C07H 19/00; C07K 15/00; C12P 21/00, 21/08

A. CLASSIFICATION OF SUBJECT MATTER: US CL :

424/130.1, 139.1, 150.1, 164.1, 184.1, 1.85.1, 242.1, 256.1; 435/69.1; 536/22.1, 23.7; 530/350, 387.1, 388.1

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